	M PTO- 11-200		F COMMERCE PATENT AND TRADEMARK OFFICE	ATTORNEY'S DOCKET NUMBER 550-266										
-		TRANSMITTAL LETTE	R TO THE UNITED STATES	U.Ş. APPLICATION NO. (If known, see 37 C.F.R. 1.5)										
DESIGNATED/ELECTED OFFICE (DO/EO/US) CONCERNING A FILING UNDER 35 U.S.C. 371 09/937716														
INTE	RNAT	TIONAL APPLICATION NO.	INTERNATIONAL FILING DATE	PRIORITY DATE CLAIMED										
		PCT/GB00/01211	30 March 2000	31 March 1999										
TITL	E OF	INVENTION												
		•	FACTOR											
APF	PLICA	NT(S) FOR DO/EO/US	MADEN et al	<u> </u>										
App	licant	herewith submits to the Unite	ed States Designated/Elected Office (DO/EO/	/US) the following items and other information:										
1.	1. X This is a FIRST submission of items concerning a filing under 35 U.S.C. 371.													
2.		This is a SECOND or SUBSEQUENT submission of items concerning a filling under 35 U.S.C. 371.												
3.	\boxtimes	This is an express request to begin national examination procedures (35 U.S.C. 371(f)). The submission must include items (5), (6), (9) and (21) indicated below.												
4.		The U.S. has been elected by the expiration of 19 months from the priority date (Article 31).												
5.	A copy of the International Application as filed (35 U.S.C. 371(c)(2)).													
	a.	is attached hereto (required only if not communicated by the International Bureau).												
	b.													
	c.	is not required, as the application was filed in the United States Receiving Office (RO/US).												
6.		An English language translation of the International Application as filed (35 U.S.C. 371(c)(2)).												
_	a. is attached hereto.													
"	b.	has been previously submitted under 35 U.S.C. 154(d)(4).												
7.		Amendments to the claims of the International Application under PCT Article 19 (35 U.S.C. 371(c)(3))												
	a.	are attached hereto (re	equired only if not communicated by the Inter	national Bureau).										
	b.	have been communicated by the International Bureau.												
	c.	have not been made; however, the time limit for making such amendments has NOT expired.												
	d.	have not been made and will not be made.												
8.		An English language translation of the amendments to the claims under PCT Article 19 (35 U.S.C. 371(c)(3)).												
9.		An oath or declaration of the inventor(s) (35 U.S.C. 371(c)(4)).												
10.		A English language translat Article 36 (35 U.S.C. 3	ion of the annexes of the International Prelim i71(c)(5)).	inary Examination Report under PCT										
	Iten	ns 11 To 20 below concern	document(s) or information included:											
11.	\boxtimes	An Information Disclosure S	Statement under 37 C.F.R. 1.97 and 1.98 w/P	TO-1449 Form, Int'l Search Report and References.										
12.		An assignment document for	or recording. A separate cover sheet in comp	oliance with 37 C.F.R. 3.28 and 3.31 is included.										
13.	\boxtimes	A FIRST preliminary amend	lment.											
14.		A SECOND or SUBSEQUE	NT preliminary amendment.											
15,		A substitute specification.												
16.		A change of power of attorney and/or address letter.												
17.	\boxtimes	Paper and computer-readable forms of the sequence listing in accordance with PCT Rule 13ter.2 and 35 U.S.C. 1.821-1.825.												
18.		A second copy of the put	olished international application under 35	5 U.S.C. 154(d)(4).										
19.		A second copy of the Englis	sh language translation of the international ap	oplication under 35 U.S.C. 154(d)(4).										
20.		Other items or information.												

410 Rec'd PCT/PTO 0 1 OCT 2001

U.S. APPLICATION NO. (IF)	APPLICATION NO (Pringer 27 F F 1 5) INTERNATIONAL APPLICATION NO PCT/GB00/01211					ATTORNEY'S DOCKET NUMBER 550-266							
21. The following fe						C	CALCULATIONS PTO USE ONLY						
BASIC NATIONAL F	EE (37 C.F.F	3. 1.492(a)	(1)-(5):			 ~	LOCEATIONS						
	Neither international preliminary examination fee (37 C.F.R. 1,482)												
nor international search fee (37 C.F.R. 1.445(a)(2)) paid to USPTO													
and International Search Report not prepared by the EPO or JPO\$1040.00													
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 a. \(\) A check in the amount of \$1188.00 to cover the above fees is enclosed. b. \(\) Please charge my Deposit Account No. 14-1140 in the amount of \$\) to cover the above fees. A duplicate copy of this form is enclosed. c. \(\) The Commissioner is hereby authorized to charge any additional fees which may be required, or credit any overpayment to Deposit Account No. 14-1140. A duplicate copy of this form is enclosed. d. \(\) The entire content of the foreign application(s), referred to in this application is/are hereby incorporated by reference in this application. 													
NOTE: Where an appropriate time limit under 37 C.F.R. 1.494 or 1.495 has not been met, a petition to revive (37 C.F.R. 1.137(a) or (b)) must be filed and granted to restore the application to pending status.													
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	October 1, 2001												
				36,663 REGISTRAT	ION NUMBE	R	Date						

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re Patent Application of

MADEN et al

Atty. Ref.: 550-266

National Phase of Int'l Appln. No. PCT/GB00/01211

Group:

(Filed: 30 March 2000)

Examiner:

For: FACTOR

October 1, 2001

Assistant Commissioner for Patents Washington, DC 20231

Sir:

PRELIMINARY AMENDMENT

In order to place the above-identified application in better condition for examination, please amend the application as follows:

IN THE SPECIFICATION

Substitute the Sequence Listing submitted herewith for that originally filed.

IN THE CLAIMS

Please substitute the following amended claims for corresponding claims previously presented. A copy of the amended claims showing current revisions is attached.

7. (Amended) A method according to claim 5, wherein said RAR β 2 receptor is administered by an entity comprising a RAR β 2 expression system.

REMARKS

The specification has been amended to make reference to include the Sequence Listing submitted herewith on separate sheets. Entry of the Sequence Listing does not raise the issue of new matter as the sequence information contained therein is presented in the application as originally filed. The computer readable copy of the Sequence Listing submitted herewith is believed to be the same as the attached paper copy of that Listing.

Attached hereto is a marked-up version of the changes made to the specification and claims by the current amendment. The attached page is captioned "Version With Markings To Show Changes Made."

Respectfully submitted,

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VERSION WITH MARKINGS TO SHOW CHANGES MADE

IN THE CLAIMS

7. (Amended) A method according to claim 5 [or claim 6], wherein said RARβ2 receptor is administered by an entity comprising a RARβ2 expression system.

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FACTOR

FIELD OF THE INVENTION

5 The present invention relates to a factor relating to neurite growth.

BACKGROUND TO THE INVENTION

It is desirable to cause neurite development, such as neurite outgrowth and/or neurite regeneration, for example in cases of nervous injuries such as spinal cord injuries.

Nerve growth factor (NGF) is known to stimulate certain events such as neurite outgrowth. However, NGF is a relatively large molecule with a correspondingly high molecular weight. Moreover, NGF is susceptible to protease mediated degradation. Due to these and other considerations, NGF is difficult to administer. NGF is also relatively expensive to prepare. These are problems associated with the prior art.

SUMMARY OF THE INVENTION

We have surprisingly found that it is possible to cause neurite development, such as neurite outgrowth and/or neurite regeneration, by using retinoic acid receptor β2 (RARβ2) and/or an agonist thereof.

SUMMARY ASPECTS OF THE PRESENT INVENTION

The present invention is based on the surprising finding that it is possible to cause neurite development, such as neurite outgrowth and/or neurite regeneration, by using RARβ2 and/or an agonist thereof.

Aspects of the present invention utilise this finding. For example it is possible to have a method that causes modulation of neurite development, such as neurite outgrowth and/or neurite regeneration, by using RARβ2 and/or an agonist thereof as explained herein.

DETAILED ASPECTS OF THE PRESENT INVENTION

In one aspect, the present invention relates to the use of RAR β 2 and/or an agonist thereof in the preparation of a medicament to cause neurite development.

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In the present invention the RAR β 2 and/or an agonist can be termed a pharmaceutically active agent.

Neurites are well known structures which develop from various neuronal cell types.

They appear as microscopic branch or comb-like structures or morphological projections from the surface of the cell from which they emanate. Examples of neurite outgrowth are shown in the accompanying figures, and in publications such as those referenced in (Maden 1998-review article), and are well known in the art.

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The RAR β 2 coding sequence (i.e. the RAR β 2 gene) is used as described hereinbelow. The RAR β 2 gene may be prepared by use of recombinant DNA techniques and/or by synthetic techniques. For example, it may be prepared using the PCR amplified gene fragment prepared as in the Examples section of this document using the primers etc. detailed therein, or it may be prepared according to any other suitable method known in the art.

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In another aspect, the present invention relates to the use of RAR β 2 and/or an agonist thereof in the preparation of a medicament to cause neurite development, wherein said agonist is retinoic acid (RA) and/or CD2019.

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Retinoic acid is commercially available. CD2019 is a polycyclic heterocarbyl molecule which is a RARβ2 agonist having the structure as discussed herein and as shown in (Elmazar *et al.*, (1996) Teratology vol. 53 pp158-167).

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In another aspect, the present invention relates to the use of RAR β 2 and/or an agonist thereof in the preparation of a medicament for the treatment of a neurological disorder.

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In another aspect, the present invention relates to the use of RAR β 2 and/or an agonist thereof in the preparation of a medicament for the treatment of a neurological disorder, wherein said neurological disorder comprises neurological injury.

In another aspect, the present invention relates to a method of treating a neurological disorder comprising administering a pharmacologically active amount of an RARβ2 receptor, and/or an agonist thereof.

In another aspect, the present invention relates to a method of treating a neurological disorder comprising administering a pharmacologically active amount of an RARβ2 receptor, and/or an agonist thereof, wherein said agonist is RA and/or CD2019.

In another aspect, the present invention relates to a method of treating a neurological disorder comprising administering a pharmacologically active amount of an RAR β 2 receptor, and/or an agonist thereof, wherein said RAR β 2 receptor is administered by an entity comprising a RAR β 2 expression system.

In another aspect, the present invention relates to a method of causing neurite development in a subject, said method comprising providing a nucleic acid construct capable of directing the expression of at least part of a RARB2 receptor, introducing said construct into one or more cells of said subject, and optionally administering a RARB2 agonist, such as RA and/or CD2019, to said subject.

In a further aspect, the invention relates to an assay method for determining whether an agent is capable of modulating RAR β 2 signalling, said method comprising providing neural cells, contacting said cells with said agent, and assessing the activity of the RAR β 2 receptor, such as through the monitoring of neurite outgrowth.

Neural cells for use in the assay method of the invention may be any suitable neural cell line, whether stably maintained in culture, or primary cells derived from an animal directly. Preferably said cells will be embryonic mouse dorsal root ganglion (DRG) cells prepared as described hereinbelow.

In a further aspect, the invention relates to a process comprising the steps of (i) performing the assay for modulation of RAR β 2 signalling described above, (ii) identifying one or more agents that are capable of modulating said RAR β 2 signalling, and (iii) preparing a quantity of those one or more identified agents.

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In a further aspect, the invention relates to a process comprising the steps of (i) performing the assay for modulation of RARβ2 signalling described above, (ii) identifying one or more agents that are capable of modulating said RARβ2 signalling, (iii) preparing a quantity of those one or more identified agents, and (iv) preparing a pharmaceutical composition comprising those one or more identified agents.

In a further aspect, the invention relates to a method of affecting the *in vivo* activity of RAR β 2 with an agent, wherein the agent is capable of modulating RAR β 2 signalling, for example capable of modulating RAR β 2 signalling in an *in vitro* assay method as described above.

In a further aspect, the invention relates to the use of an agent in the preparation of a pharmaceutical composition for the treatment of a neurological disorder or injury, wherein the agent is capable of modulating RAR β 2 signalling, for example capable of modulating RAR β 2 signalling in an *in vitro* assay method as described above.

In a further aspect, the invention relates to a method of treating a subject with an agent, wherein the agent is capable of modulating RAR β 2 signalling for example capable of modulating RAR β 2 signalling in an *in vitro* assay method as described above.

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In a further aspect, the invention relates to a pharmaceutical composition comprising RARβ2 and/or an agonist thereof in admixture with a pharmaceutically acceptable carrier, diluent or excipient; wherein the pharmaceutical composition is for use to cause neurite development.

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For ease of reference, these and further aspects of the present invention are now discussed under appropriate section headings. However, the teachings under each section are not necessarily limited to each particular section.

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PREFERABLE ASPECTS

In a preferred aspect, the administration of a nucleic acid construct capable of directing the expression of RAR β 2 will be accompanied by the administration of a RAR β 2 agonist such as RA, or preferably CD2019 (or a mimetic thereof).

Preferably said agonist will be to some degree selective for the RAR β 2 receptor. Preferably said agonist will not significantly affect the RAR α receptor. Preferably said agonist will not significantly affect the RAR α receptor. More preferably said agonist will not significantly affect the RAR α receptor or the RAR α receptor. Even more preferably, said agonist will exhibit a high degree of selectivity for the RAR α 2 receptor.

In a preferred aspect, the administration of a nucleic acid construct capable of directing the expression of RARβ2 will be accomplished using a vector, preferably a viral vector, more preferably a retroviral vector. In a highly preferred embodiment, the administration of a nucleic acid construct capable of directing the expression of RARβ2 will be accomplished using a retroviral vector capable of infecting non-dividing mammalian cells such as neural cells.

ADVANTAGES

The present invention is advantageous because RARβ2 and/or an agonist thereof can cause modulation of neural cell development.

It is also an advantage of the present invention that administration of NGF to a subject is avoided.

30 It is also an advantage of the present invention that it enable neurite outgrowth to be promoted in adult neural tissue.

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RETINOIDS

Retinoids are a family of molecules derived from vitamin A and include the biologically active metabolite, retinoic acid (RA). The cellular effects of RA are mediated through the action of two classes of receptors, the retinoic acid receptors (RARs) which are activated both by alltrans-RA (tRA) and 9-cis-RA (9-cis-RA), and the retinoid X receptors (RXRs), which are activated only by 9-cis-RA (Kastner et al., 1994; Kleiwer et al., 1994). The receptors are of three major subtypes, α , β and γ , of which there are multiple isoforms due to alternative splicing and differential promoter usage (Leid et al.). The RARs mediate gene expression by forming heterodimers with the RXRs, whilst the RXRs can mediate gene expression as homodimers or by forming heterodimers with a variety of orphan receptors (Mangelsdorf & Evans, 1995). Many studies on a variety of embryonic neuronal types have shown that RA can stimulate both neurite number and length (review, Maden, 1998), as, indeed, can the neurotrophins (Campenot, 1977; Lindsay, 1988; Tuttle and Mathew, 1995). The neurotrophins are a family of growth factors that are required for the survival of a variety of neurons of primary sensory neurons in the developing peripheral nervous system (Snider, 1994). One of the earliest genes induced by NGF in PC12 cells is the orphan receptor NGFI-B (NURR1) (Millbrandt, 1989). This suggests that the growth factor and retinoid mediated pathway in developing neurons can interact.

Background teachings on these aspects have been presented by Victor A. McKusick et al on http://www.ncbi.nlm.nih.gov/Omim. The following information has been extracted from that source.

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Three retinoic acid receptors, alpha, beta, and gamma, are members of the nuclear receptor superfamily. Retinoic acid was the first morphogen described in vertebrates. The RARA and RARB genes are more homologous to those of the 2 closely related thyroid hormone receptors THRA and THRB, located on chromosomes 17 and 3, respectively, than to any other members of the nuclear receptor family. These observations suggest that the thyroid hormone and retinoic acid receptors evolved by gene, and possibly chromosome, duplications from a common ancestor which itself diverged rather early in evolution from the common ancestor of the steroid receptor group of the family. The RARB gene, formerly symbolized HAP, maps to 3p24 by somatic cell hybridization and in situ hybridization.

Benbrook et al. (1988) showed a predominant distribution in epithelial tissues and therefore used the designation RAR(epsilon). By in situ hybridization, Mattei et al. (1988) assigned the RARB gene to 3p24. Using deletion mapping, de The et al. (1990) identified a 27-bp fragment located 59-bp upstream

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of the transcriptional start, which confers retinoic acid responsiveness on the herpesvirus thymidine kinase promoter. They found indications that both alpha and beta receptors act through the same DNA sequence. Matter et al. (1991) assigned the corresponding gene to chromosome 14, band A, in the mouse, and to chromosome 15 in the rat.

- Nadeau et al. (1992) confirmed assignment of the mouse homolog to the centromeric portion of chromosome 14.
 - From a comparison of a hepatitis-B virus (HBV) integration site present in a particular human hepatocellular carcinoma (HCC) with the corresponding unoccupied site in the nontumorous tissue of the same liver, Dejean et al. (1986) found that HBV integration placed the viral sequence next to a liver cell sequence that bears a striking resemblance to both an oncogene, ERBA, and the supposed DNA-binding domain of the human glucocorticoid receptor and estrogen receptor genes.
 - Dejean et al. (1986) suggested that this gene, usually silent or transcribed at a very low level in normal hepatocytes, becomes inappropriately expressed as a consequence of HBV integration, thus contributing to the cell transformation.
- By means of a panel of rodent-human somatic cell hybrid DNAs, Dejean et al. (1986) localized the gene to chromosome 3. Further studies by de The et al. (1987) suggested that the HAP gene product may be a novel ligand-responsive regulatory protein whose inappropriate expression in liver is related to hepatocellular carcinogenesis. Brand et al. (1988) showed that the novel protein called HAP (for HBV-activated protein) is a retinoic acid receptor. They referred to this receptor as the beta type (RARB) and mapped it to 3p25-p21.
 - Lotan et al. (1995) found that the expression of RARB mRNA is selectively lost in premalignant oral lesions and can be restored by treatment with isotretinoin. Restoration of the expression of RARB mRNA was associated with a clinical response.
- RARB, RARG, RXRB, and RXRG are expressed in the striatum. To study the effect of these genes on locomotion, Kreczel et al. (1998) developed single and double knockout mice and analyzed their 25 locomotor skills by open field and rotarod testing. RARB-RXRB, RARB-RXRG, and RXRB-RXRG double null mutant mice, but not the corresponding single null mutants, exhibited reductions in forward locomotion when compared with wildtype littermates. Forty percent of the RARB-RXRB null mutants showed backward locomotion. Rotarod test performance was impaired for RARB, RARB-RXRB, RARB-RXRG, and RXRB-RXRG mice. In contrast, RARA, RARG, RARA-RXRG, and RARG-30 RXRG null mice showed no defects in locomotion, even though both RARA and RARG are also expressed in the striatum. The morphology, development, and function of skeletal muscle, peripheral nerves, and spinal cord were normal in all single and double null mutants, as were balance reflexes. These results suggested to Kreczel et al. (1998) that RARB, RXRB, and RXRG are involved specifically in the control of locomotor behaviors, and that heterodimers of RARB with either RXRB or 35 RXRG are the functional receptor units, such that RXRB and RXRG are functionally redundant.
 - Kreczel et al. (1998)studied the expression of D1 and D2 dopamine receptors (D1R and D2R), the most abundant dopamine receptors in the striatum, in these mutant mice. RARB-RXRB, RARB-RXRG, and

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RXRB-RXRG double null mutants, but not RARB or RXRG single mutants, exhibited 40% and 30% reduction in whole-striatal D1R and D2R transcripts, respectively, when compared with wildtype controls.

The reduction was mostly in the medioventral regions of the striatum, including the shell and core of the nucleus accumbens, and the mediodorsal part of the caudate putamen. The reduction was not due to loss of D2R-expressing neurons; no increase in apoptosis was noted. The histology of the striatum was normal.

The characterization of a retinoic acid response element in the D2R promoter by Samad et al. (1997) led Kreczel et al. (1998) to suggest that the reduction in D2R and D2R expression occurs on a transcriptional level. The RARB-RXRB, RARB-RXRG, and RXRB-RXRG double null mutants did not exhibit the normal increase in locomotion induced by cocaine, mimicking the phenotype of D1R-null mice.

Taken together, these results indicated to Kreczel et al. (1998) that retinoids are involved in controlling the function of the dopaminergic mesolimbic pathway and suggested that defects in retinoic acid signaling may contribute to neurological disorders.

AGONISTS

The agonist of the present invention may be any suitable RAR β 2 agonist. Preferably, said agonist of RAR β 2 is capable of activating RAR β 2 in a transactivation assay.

The agonist may be an organic compound or other chemical. The agonist can be an amino acid sequence or a chemical derivative thereof, or a combination thereof. The agent may even be a nucleotide sequence - which may be a sense sequence or an anti-sense sequence. The agent may even be an antibody.

Typically, the agonist will be an organic compound. Typically the organic compound will comprise two or more hydrocarbyl groups. Here, the term "hydrocarbyl group" means a group comprising at least C and H and may optionally comprise one or more other suitable substituents. Examples of such substituents may include halo-, alkoxy-, nitro-, an alkyl group, a cyclic group etc. In addition to the possibility of the substituents being a cyclic group, a combination of substituents may form a cyclic group. If the hydrocarbyl group comprises more than one C then those carbons need not necessarily be linked to each other. For example, at least two of the carbons may be linked *via* a suitable element or group. Thus, the hydrocarbyl group may contain hetero atoms. Suitable hetero atoms will be apparent to those skilled in the art and

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include, for instance, sulphur, nitrogen and oxygen. For some applications, preferably the agent comprises at least one cyclic group. The cyclic group may be a polycyclic group, such as a non-fused polycyclic group. For some applications, the agonist comprises at least the one of said cyclic groups linked to another hydrocarbyl group.

SPECIFIC AGONISTS

An example of a specific agonist according to the present invention is retinoic acid (RA). Both common forms of retinoic acid (either all-trans retinoic acid (tRA), or 9-cis-RA) are agonists of RARβ2.

CD2019 is a RARβ2 agonist having the structure as discussed herein and as shown in (Elmazar *et al.*, (1996) Teratology vol. 53 pp158-167). This and other agonists are also discussed in (Beard and Chandraratna p.194; Johnson *et al.*, 1996). The structure of CD2019 is presented as Formula I in the attached figures.

An alternative RARβ2 agonist is presented as Formula II in the attached figures.

The present invention also encompasses mimetics or bioisosteres of the formulae of Formula I and/or Formula II.

Preferably the agonist useful according to the present invention is selective for RARβ2.

25 ASSAY TO DETERMINE RARβ2 AGONISM

Examples of agonists according to the present invention may be identified and/or verified by using an assay to determine RARβ2 agonism.

Hence, the present invention also encompasses (i) determining if a candidate agent is capable of acting as a RARβ2 agonist; (ii) if said candidate agent is capable of acting as a RARβ2 agonist then delivering said agent to a subject and in such an amount to cause neurite development.

ASSAY

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Any one or more of appropriate targets - such as an amino acid sequence and/or nucleotide sequence - may be used for identifying an agent capable of modulating RARβ2 in any of a variety of drug screening techniques. The target employed in such a test may be free in solution, affixed to a solid support, borne on a cell surface, or located intracellularly. The abolition of target activity or the formation of binding complexes between the target and the agent being tested may be measured.

The assay of the present invention may be a screen, whereby a number of agents are tested. In one aspect, the assay method of the present invention is a high through put screen.

Techniques for drug screening may be based on the method described in Geysen,
European Patent Application 84/03564, published on September 13, 1984. In
summary, large numbers of different small peptide test compounds are synthesized
on a solid substrate, such as plastic pins or some other surface. The peptide test
compounds are reacted with a suitable target or fragment thereof and washed.
Bound entities are then detected - such as by appropriately adapting methods well
known in the art. A purified target can also be coated directly onto plates for use in a
drug screening techniques. Alternatively, non-neutralising antibodies can be used to
capture the peptide and immobilise it on a solid support.

This invention also contemplates the use of competitive drug screening assays in which neutralising antibodies capable of binding a target specifically compete with a test compound for binding to a target.

Another technique for screening provides for high throughput screening (HTS) of agents having suitable binding affinity to the substances and is based upon the method described in detail in WO-A-84/03564.

It is expected that the assay methods of the present invention will be suitable for both small and large-scale screening of test compounds as well as in quantitative assays.

35 In one preferred aspect, the present invention relates to a method of identifying

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agents that selectively modulate RARβ2.

In a preferred aspect, the assay of the present invention utilises cells that display RAR β 2 on their surface. These cells may be isolated from a subject possessing such cells. However, preferably, the cells are prepared by transfecting cells so that upon transfect those cells display on their surface RAR β 2.

Another example of an assay that may be used is described in WO-A-9849271, which concerns an immortalised human terato-carcinoma CNS neuronal cell line, which is siad to have a high level of neuronal differentiation and is useful in detecting compounds which bind to RAR62.

REPORTERS

A wide variety of reporters may be used in the assay methods (as well as screens) of the present invention with preferred reporters providing conveniently detectable signals (eg. by spectroscopy). By way of example, a reporter gene may encode an enzyme which catalyses a reaction which alters light absorption properties.

Other protocols include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA) and fluorescent activated cell sorting (FACS). A two-site, monoclonal-based immunoassay utilising monoclonal antibodies reactive to two non-interfering epitopes may even be used. These and other assays are described, among other places, in Hampton R et al (1990, Serological Methods, A Laboratory Manual, APS Press, St Paul MN) and Maddox DE et al (1983, J Exp Med 15 8:121 1).

Examples of reporter molecules include but are not limited to (galactosidase, invertase, green fluorescent protein, luciferase, chloramphenicol, acetyltransferase, (glucuronidase, exo-glucanase and glucoamylase. Alternatively, radiolabelled or fluorescent tag-labelled nucleotides can be incorporated into nascent transcripts which are then identified when bound to oligonucleotide probes.

By way of further examples, a number of companies such as Pharmacia Biotech (Piscataway, NJ), Promega (Madison, WI), and US Biochemical Corp (Cleveland, OH) supply commercial kits and protocols for assay procedures. Suitable reporter

molecules or labels include those radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents as well as substrates, cofactors, inhibitors, magnetic particles and the like. Patents teaching the use of such labels include US-A-3817837; US-A-3850752: US-A-3939350; US-A-3996345; US-A-4277437; US-A-4275149 and US-A-4366241.

HOST CELLS

Polynucleotides for use in the present invention – such as for use as targets or for expressing targets or for use as the pharmaceutically active agent - may be introduced into host cells.

The term "host cell" - in relation to the present invention includes any cell that could comprise the polynucleotide sequence of the present invention.

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Here, polynucleotides may be introduced into prokaryotic cells or eukaryotic cells, for example yeast, insect or mammalian cells.

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Polynucleotides of the invention may introduced into suitable host cells using a variety of techniques known in the art, such as transfection, transformation and electroporation. Where polynucleotides of the invention are to be administered to animals, several techniques are known in the art, for example infection with recombinant viral vectors such as retroviruses, herpes simplex viruses and adenoviruses, direct injection of nucleic acids and biolistic transformation.

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Thus, a further embodiment of the present invention provides host cells transformed or transfected with a polynucleotide that is or expresses the target of the present invention. Preferably said polynucleotide is carried in a vector for the replication and expression of polynucleotides that are to be the target or are to express the target. The cells will be chosen to be compatible with the said vector and may for example be prokaryotic (for example bacterial), fungal, yeast or plant cells.

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The gram negative bacterium *E. coli* is widely used as a host for heterologous gene expression. However, large amounts of heterologous protein tend to accumulate inside the cell. Subsequent purification of the desired protein from the bulk of E.coli

intracellular proteins can sometimes be difficult.

In contrast to *E.coli*, bacteria from the genus *Bacillus* are very suitable as heterologous hosts because of their capability to secrete proteins into the culture medium. Other bacteria suitable as hosts are those from the genera Streptomyces and Pseudomonas.

Depending on the nature of the polynucleotide encoding the polypeptide of the present invention, and/or the desirability for further processing of the expressed protein, eukaryotic hosts such as yeasts or other fungi may be preferred. In general, yeast cells are preferred over fungal cells because they are easier to manipulate. However, some proteins are either poorly secreted from the yeast cell, or in some cases are not processed properly (e.g. hyperglycosylation in yeast). In these instances, a different fungal host organism should be selected.

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Examples of suitable expression hosts within the scope of the present invention are fungi such as Aspergillus species (such as those described in EP-A-0184438 and EP-A-0284603) and Trichoderma species; bacteria such as Bacillus species (such as those described in EP-A-0134048 and EP-A-0253455), Streptomyces species and Pseudomonas species; and yeasts such as Kluyveromyces species (such as those described in EP-A-0096430 and EP-A-0301670) and Saccharomyces species. By way of example, typical expression hosts may be selected from Aspergillus niger, Aspergillus niger var. tubigenis, Aspergillus niger var. awamori, Aspergillus aculeatis, Aspergillus nidulans, Aspergillus orvzae, Trichoderma reesei, Bacillus subtilis, Bacillus licheniformis, Bacillus amyloliquefaciens, Kluyveromyces lactis and Saccharomyces cerevisiae.

Polypeptides that are extensively modified may require correct processing to complete their function. In those instances, mammalian cell expression systems (such as HEK-293, CHO, HeLA) are required, and the polypeptides are expressed either intracellularly, on the cell membranes, or secreted in the culture media if preceded by an appropriate leader sequence.

The use of suitable host cells - such as yeast, fungal, plant and mammalian host cells - may provide for post-translational modifications (e.g. myristoylation, glycosylation,

truncation, lipidation and tyrosine, serine or threonine phosphorylation) as may be needed to confer optimal biological activity on recombinant expression products of the present invention.

5 ORGANISM

The term "organism" in relation to the present invention includes any organism that could comprise the sequence according to the present invention and/or products obtained therefrom. Examples of organisms may include a fungus, yeast or a plant.

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The term "transgenic organism" in relation to the present invention includes any organism that comprises the target according to the present invention and/or products obtained.

15 TRANSFORMATION OF HOST CELLS/HOST ORGANISMS

As indicated earlier, the host organism can be a prokaryotic or a eukaryotic organism. Examples of suitable prokaryotic hosts include *E. coli* and *Bacillus subtilis*. Teachings on the transformation of prokaryotic hosts is well documented in the art, for example see Sambrook *et al* (Molecular Cloning: A Laboratory Manual, 2nd edition, 1989, Cold Spring Harbor Laboratory Press) and Ausubel et al., Current Protocols in Molecular Biology (1995), John Wiley & Sons, Inc.

.If a prokaryotic host is used then the nucleotide sequence may need to be suitably modified before transformation - such as by removal of introns.

In another embodiment the transgenic organism can be a yeast. In this regard, yeast have also been widely used as a vehicle for heterologous gene expression. The species Saccharomyces cerevisiae has a long history of industrial use, including its use for heterologous gene expression. Expression of heterologous genes in Saccharomyces cerevisiae has been reviewed by Goodey et al (1987, Yeast Biotechnology, D R Berry et al, eds, pp 401-429, Allen and Unwin, London) and by King et al (1989, Molecular and Cell Biology of Yeasts, E F Walton and G T Yarronton, eds, pp 107-133, Blackie, Glasgow).

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For several reasons Saccharomyces cerevisiae is well suited for heterologous gene expression. First, it is non-pathogenic to humans and it is incapable of producing certain endotoxins. Second, it has a long history of safe use following centuries of commercial exploitation for various purposes. This has led to wide public acceptability. Third, the extensive commercial use and research devoted to the organism has resulted in a wealth of knowledge about the genetics and physiology as well as large-scale fermentation characteristics of Saccharomyces cerevisiae.

A review of the principles of heterologous gene expression in *Saccharomyces* cerevisiae and secretion of gene products is given by E Hinchcliffe E Kenny (1993, "Yeast as a vehicle for the expression of heterologous genes", Yeasts, Vol 5, Anthony H Rose and J Stuart Harrison, eds, 2nd edition, Academic Press Ltd.).

Several types of yeast vectors are available, including integrative vectors, which require recombination with the host genome for their maintenance, and autonomously replicating plasmid vectors.

In order to prepare the transgenic Saccharomyces, expression constructs are prepared by inserting the nucleotide sequence of the present invention into a construct designed for expression in yeast. Several types of constructs used for heterologous expression have been developed. The constructs contain a promoter active in yeast fused to the nucleotide sequence of the present invention, usually a promoter of yeast origin, such as the GAL1 promoter, is used. Usually a signal sequence of yeast origin, such as the sequence encoding the SUC2 signal peptide, is used. A terminator active in yeast ends the expression system.

For the transformation of yeast several transformation protocols have been developed. For example, a transgenic Saccharomyces according to the present invention can be prepared by following the teachings of Hinnen et al (1978, Proceedings of the National Academy of Sciences of the USA 75, 1929); Beggs, J D (1978, Nature, London, 275, 104); and Ito, H et al (1983, J Bacteriology 153, 163-168).

The transformed yeast cells are selected using various selective markers. Among the markers used for transformation are a number of auxotrophic markers such as LEU2,

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HIS4 and TRP1, and dominant antibiotic resistance markers such as aminoglycoside antibiotic markers, eq G418.

Another host organism is a plant. The basic principle in the construction of genetically modified plants is to insert genetic information in the plant genome so as to obtain a stable maintenance of the inserted genetic material. Several techniques exist for inserting the genetic information, the two main principles being direct introduction of the genetic information and introduction of the genetic information by use of a vector system. A review of the general techniques may be found in articles by Potrykus (Annu Rev Plant Physiol Plant Mol Biol [1991] 42:205-225) and Christou (Agro-Food-Industry Hi-Tech March/April 1994 17-27). Further teachings on plant transformation may be found in EP-A-0449375.

Further hosts suitable for the nucleotide sequence of the present invention include higher eukaryotic cells, such as insect cells or vertebrate cells, particularly mammalian cells, including human cells, or nucleated cells from other multicellular organisms. In recent years propagation of vertebrate cells in culture (tissue culture) has become a routine procedure. Examples of useful mammalian host cell lines are epithelial or fibroblastic cell lines such as Chinese hamster ovary (CHO) cells, NIH 3T3 cells, HeLa cells or 293T cells.

The nucleotide sequence of the present invention may be stably incorporated into host cells or may be transiently expressed using methods known in the art. By way of example, stably transfected mammalian cells may be prepared by transfecting cells with an expression vector having a selectable marker gene, and growing the transfected cells under conditions selective for cells expressing the marker gene. To prepare transient transfectants, mammalian cells are transfected with a reporter gene to monitor transfection efficiency.

To produce such stably or transiently transfected cells, the cells should be transfected with a sufficient amount of the nucleotide sequence of the present invention. The precise amounts of the nucleotide sequence of the present invention may be empirically determined and optimised for a particular cell and assay.

35 Thus, the present invention also provides a method of transforming a host cell with a

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nucleotide sequence that is to be the target or is to express the target. Host cells transformed with the nucleotide sequence may be cultured under conditions suitable for the expression of the encoded protein. The protein produced by a recombinant cell may be displayed on the surface of the cell. If desired, and as will be understood by those of skill in the art, expression vectors containing coding sequences can be designed with signal sequences which direct secretion of the coding sequences through a particular prokaryotic or eukaryotic cell membrane. Other recombinant constructions may join the coding sequence to nucleotide sequence encoding a polypeptide domain which will facilitate purification of soluble proteins (Kroll DJ et al (1993) DNA Cell Biol 12:441-53).

RECEPTORS

The RARβ2 receptor as discussed herein includes mimetics, homologues, fragments and part or all of the entire gene product. Preferably the RARβ2 receptor as discussed herein refers to substantially the entire gene product.

NEUROLOGICAL DISORDERS

The term neurological disorders as used herein may refer to any injury, whether mechanically (for example by trauma) or chemically induced (for example by neurotoxin(s), or by an regime of treatment having an immunosuppressant effect, whether by design, or as a side-effect), any neural pathology such as caused by viral infection or otherwise, any degenerative disorder, or other nerve tissue related disorder.

Examples of neurological disorders include conditions such as Parkinson's disease, Alzheimer's disease, senility, motor neurone disease, schizophrenia as well as other neural and/or neurodegenerative disorders. Other neural related disorders may include glaucoma or other cause of damage to the optic nerve, Bell's palsy or other forms of localised paralysis, neurally based impotence such as caused by nerve trauma following radical prostatectomy, or other complaints. Other disorders in which the invention may be useful include neuropathological effects of diabetes, AIDS neuropathy, leprosy etc.

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The term neurological disorder refers to any disorder of a nervous system, whether the peripheral nervous system or the central nervous system (CNS), whether the sympathetic nervous system, or the parasympathetic nervous system, or whether affecting a subset or superset of different nerve types.

5 NUCLEOTIDE OF INTEREST (NOI)

In accordance with the present invention, the NOI sequence may encode a peptide which peptide may be the pharmaceutically active agent – such as an RA receptor, preferably RARβ2, or an agonist thereof.

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Such coding NOI sequences may be typically operatively linked to a suitable promoter capable of driving expression of the peptide, such as in one or more specific cell types.

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In addition to the NOI or part thereof and the expression regulatory elements described herein, the delivery system may contain additional genetic elements for the efficient or regulated expression of the gene or genes, including promoters/enhancers, translation initiation signals, internal ribosome entry sites (IRES), splicing and polyadenylation signals.

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The NOI or NOIs may be under the expression control of an expression regulatory element, usually a promoter or a promoter and enhancer. The enhancer and/or promoter may be preferentially active in neural cells, such that the NOI is preferentially expressed in the particular cells of interest, such as in nerve cells. Thus any significant biological effect or deleterious effect of the NOI on the individual being treated may be reduced or eliminated. The enhancer element or other elements conferring regulated expression may be present in multiple copies. Likewise, or in addition, the enhancer and/or promoter may be preferentially active in one or more specific cell types - such as neural cells for example post-mitotically terminally differentiated non-replicating cells such as neurons.

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The term "promoter" is used in the normal sense of the art, e.g. an RNA polymerase binding site in the Jacob-Monod theory of gene expression.

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The term "enhancer" includes a DNA sequence which binds to other protein components of the transcription initiation complex and thus facilitates the initiation of transcription directed by its associated promoter.

5 EXPRESSION VECTOR

Preferably, the NOI (e.g. that encoding RARβ2 or part thereof) used in the method of the present invention is inserted into a vector which is operably linked to a control sequence that is capable of providing for the expression of the coding sequence by the host cell, i.e. the vector is an expression vector.

TARGETED VECTOR

The term "targeted vector" refers to a vector whose ability to infect/transfect/transduce
a cell or to be expressed in a host and/or target cell is restricted to certain cell types
within the host organism, usually cells having a common or similar phenotype.

DELIVERY

- The delivery system for use in the present invention may be any suitable delivery system for delivering said NOI and providing said NOI is expressed *in vivo* to produce said associated peptide (e.g. RARβ2), which in turn provides the beneficial therapeutic effect.
- The delivery system may be a viral delivery system. Viral delivery systems include but are not limited to adenovirus vector, an adeno-associated viral (AAV) vector, a herpes viral vector, retroviral vector, lentiviral vector, baculoviral vector. Alternatively, the delivery system may be a non-viral delivery system such as by way of example DNA transfection methods of, for example, plasmids, chromosomes or artificial chromosomes. Here transfection includes a process using a non-viral vector to deliver a gene to a target mammalian cell. Typical transfection methods include electroporation, DNA biolistics, lipid-mediated transfection, compacted DNA-mediated transfection, liposomes, immunoliposomes, lipofectin, cationic agent-mediated, cationic facial amphiphiles (CFAs) (Nature Biotechnology 1996 14; 556), and combinations thereof.

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Other examples of vectors include *ex vivo* delivery systems - which include but are not limited to DNA transfection methods such as electroporation, DNA biolistics, lipid-mediated transfection, compacted DNA-mediated transfection).

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In a preferred aspect, the delivery system is a vector.

In a more preferred aspect, the delivery system is a viral delivery system - sometimes referred to as a viral vector.

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VECTORS

As it is well known in the art, a vector is a tool that allows or faciliates the transfer of an entity from one environment to another. By way of example, some vectors used in recombinant DNA techniques allow entities, such as a segment of DNA (such as a heterologous DNA segment, such as a heterologous cDNA segment), to be transferred into a target cell. Optionally, once within the target cell, the vector may then serve to maintain the heterologous DNA within the cell or may act as a unit of DNA replication. Examples of vectors used in recombinant DNA techniques include plasmids, chromosomes, artificial chromosomes or viruses.

The term "vector" includes expression vectors and/or transformation vectors.

.The term "expression vector" means a construct capable of *in vivo* or *in vitrolex vivo* expression.

The term "transformation vector" means a construct capable of being transferred from one species to another.

30 VIRAL VECTORS

In the present invention, the NOI may be introduced into suitable host cells using a viral delivery system (a viral vector). A variety of viral techniques are known in the art, such as for example infection with recombinant viral vectors such as retroviruses, herpes simplex viruses and adenoviruses.

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Suitable recombinant viral vectors include but are not limited to adenovirus vectors, adeno-associated viral (AAV) vectors, herpes-virus vectors, a retroviral vector, lentiviral vectors, baculoviral vectors, pox viral vectors or parvovirus vectors (see Kestler et al 1999 Human Gene Ther 10(10):1619-32). In the case of viral vectors, gene delivery is typically mediated by viral infection of a target cell.

RETROVIRAL VECTORS

Examples of retroviruses include but are not limited to: murine leukemia virus (MLV), human immunodeficiency virus (HIV), equine infectious anaemia virus (EIAV), mouse mammary tumour virus (MMTV), Rous sarcoma virus (RSV), Fujinami sarcoma virus (FuSV), Moloney murine leukemia virus (Mo-MLV), FBR murine osteosarcoma virus (FBR MSV), Moloney murine sarcoma virus (Mo-MSV), Abelson murine leukemia virus (A-MLV), Avian myelocytomatosis virus-29 (MC29), and Avian erythroblastosis virus (AEV).

Preferred vectors for use in accordance with the present invention are recombinant viral vectors, in particular recombinant retroviral vectors (RRV) such as lentiviral vectors.

The term "recombinant retroviral vector" (RRV) refers to a vector with sufficient retroviral genetic information to allow packaging of an RNA genome, in the presence of packaging components, into a viral particle capable of infecting a target cell. Infection of the target cell includes reverse transcription and integration into the target cell genome. The RRV carries non-viral coding sequences which are to be delivered by the vector to the target cell. An RRV is incapable of independent replication to produce infectious retroviral particles within the final target cell. Usually the RRV lacks a functional *gag-pol* and/or *env* gene and/or other genes essential for replication.

A detailed list of retroviruses may be found in Coffin *et al* ("Retroviruses" 1997 Cold Spring Harbour Laboratory Press Eds: JM Coffin, SM Hughes, HE Varmus pp 758-763).

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NON-VIRAL DELIVERY

The pharmaceutically active agent (e.g. the RARβ2) may be administered using non-viral techniques.

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By way of example, the pharmaceutically active agent may be delivered using peptide delivery. Peptide delivery uses domains or sequences from proteins capable of translocation through the plasma and/or nuclear membrane

- Polypeptides of interest such as RARβ2 may be directly introduced to the cell by microinjection, or delivery using vesicles such as liposomes which are capable of fusing with the cell membrane. Viral fusogenic peptides may also be used to promote membrane fusion and delivery to the cytoplasm of the cell.
- Preferably, the RARβ2 or fragment(s) thereof may be delivered into cells as protein fusions or conjugates with a protein capable of crossing the plasma membrane and/or the nuclear membrane. Preferably, the RARβ2 or fragment(s) thereof is fused or conjugated to a domain or sequence from such a protein responsible for the translocational activity. Preferred translocation domains and sequences include domains and sequences from the HIV-1-trans-activating protein (Tat), *Drosophila* Antennapedia homeodomain protein and the herpes simplex-1 virus VP22 protein.
 - Exogenously added HIV-1-trans-activating protein (Tat) can translocate through the plasma membrane and to reach the nucleus to transactivate the viral genome. Translocational activity has been identified in amino acids 37-72 (Fawell et al., 1994, *Proc. Natl. Acad. Sci. U. S. A.* 91, 664-668), 37-62 (Anderson et al., 1993, *Biochem. Biophys. Res. Commun.* 194, 876-884) and 49-58 (having the basic sequence RKKRRQRRR) of HIV-Tat. Vives et al. (1997), *J Biol Chem* 272, 16010-7 identified a sequence consisting of amino acids 48-60 (CGRKKRRQRRRPPQC), which appears to be important for translocation, nuclear localisation and trans-activation of cellular genes. The third helix of the *Drosophila* Antennapedia homeodomain protein has also been shown to possess similar properties (reviewed in Prochiantz, A., 1999, *Ann N Y Acad Sci*, 886, 172-9). The domain responsible for translocation in Antennapedia has been localised to a 16 amino acid long peptide rich in basic amino acids having the

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sequence RQIKIWFQNRRMKWKK (Derossi, et al., 1994, *J Biol Chem*, 269, 10444-50). This peptide has been used to direct biologically active substances to the cytoplasm and nucleus of cells in culture (Theodore, et al., 1995, *J. Neurosci* 15, 7158-7167). The VP22 tegument protein of herpes simplex virus is capable of intercellular transport, in which VP22 protein expressed in a subpopulation of cells spreads to other cells in the population (Elliot and O'Hare, 1997, *Cell* 88, 223-33). Fusion proteins consisting of GFP (Elliott and O'Hare, 1999, *Gene Ther* 6, 149-51), thymidine kinase protein (Dilber et al., 1999, *Gene Ther* 6, 12-21) or p53 (Phelan et al., 1998, *Nat Biotechnol* 16, 440-3) with VP22 have been targeted to cells in this manner. Any of the domains or sequences as set out above may be used to direct RARβ2 or fragment(s) thereof into cell(s). Any of the domains or sequences as set out above, or others identified as having translocational activity, may be used to direct the RARβ2 or fragment(s) thereof into a cell.

15 PHARMACEUTICAL COMPOSITIONS

The present invention also provides a pharmaceutical composition comprising administering a therapeutically effective amount of the agent of the present invention (such as RARβ2 and/or an agonist thereof as discussed herein) and a pharmaceutically acceptable carrier, diluent or excipients (including combinations thereof).

The pharmaceutical composition may comprise two components – wherein a first component comprises RARβ2 and a second component which comprises the agonist thereof. The first and second component may be delivered sequentially, simultaneously or together, and even by different administration routes.

The pharmaceutical compositions may be for human or animal usage in human and veterinary medicine and will typically comprise any one or more of a pharmaceutically acceptable diluent, carrier, or excipient. Acceptable carriers or diluents for therapeutic use are well known in the pharmaceutical art, and are described, for example, in Remington's Pharmaceutical Sciences, Mack Publishing Co. (A. R. Gennaro edit. 1985). The choice of pharmaceutical carrier, excipient or diluent can be selected with regard to the intended route of administration and standard pharmaceutical practice.

The pharmaceutical compositions may comprise as - or in addition to - the carrier, excipient or diluent any suitable binder(s), lubricant(s), suspending agent(s), coating agent(s), solubilising agent(s).

5 Preservatives, stabilizers, dyes and even flavoring agents may be provided in the pharmaceutical composition. Examples of preservatives include sodium benzoate, sorbic acid and esters of p-hydroxybenzoic acid. Antioxidants and suspending agents may be also used.

There may be different composition/formulation requirements dependent on the different delivery systems. By way of example, the pharmaceutical composition of the present invention may be formulated to be delivered using a mini-pump or by a mucosal route, for example, as a nasal spray or aerosol for inhalation or ingestable solution, or parenterally in which the composition is formulated by an injectable form, for delivery, by, for example, an intravenous, intramuscular or subcutaneous route. Alternatively, the formulation may be designed to be delivered by both routes.

Where the agent is to be delivered mucosally through the gastrointestinal mucosa, it should be able to remain stable during transit though the gastrointestinal tract; for example, it should be resistant to proteolytic degradation, stable at acid pH and resistant to the detergent effects of bile.

Where appropriate, the pharmaceutical compositions can be administered by inhalation, in the form of a suppository or pessary, topically in the form of a lotion, solution, cream, ointment or dusting powder, by use of a skin patch, orally in the form of tablets containing excipients such as starch or lactose, or in capsules or ovules either alone or in admixture with excipients, or in the form of elixirs, solutions or suspensions containing flavouring or colouring agents, or they can be injected parenterally, for example intravenously, intramuscularly or subcutaneously. For parenteral administration, the compositions may be best used in the form of a sterile aqueous solution which may contain other substances, for example enough salts or monosaccharides to make the solution isotonic with blood. For buccal or sublingual administration the compositions may be administered in the form of tablets or lozenges which can be formulated in a conventional manner.

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PHARMACEUTICAL COMBINATIONS

The agent of the present invention may be administered with one or more other pharmaceutically active substances. By way of example, the present invention covers the simultaneous, or sequential treatments with an agent according to the present invention and one or more steroids, analgesics, antivirals or other pharmaceutically active substance(s).

It will be understood that these regimes include the administration of the substances sequentially, simultaneously or together.

EXAMPLES

The present invention will now be described, by way of example only, in which reference will be made to the following figures:

Figure 1 (which is Figure 1 referred to in Example 1) shows a photograph.

Figure 2, (which is Figure 2 referred to in Example 1) shows barcharts and a photograph.

- Figure 3, (which is Figure 1 referred to in Example 2) shows a photograph.
 - Figure 4, (which is Figure 2 referred to in Example 2) shows a photograph.
 - Figure 5, (which is Figure 3 referred to in Example 2) shows a photograph.
 - Figure 6, (which is Figure 4 referred to in Example 2) shows a photograph.
 - Figure 7, (which is Figure 5 referred to in Example 2) shows a photograph.
- Figure 8, (which is Figure 6 referred to in Example 2) shows a barchart.
 - Figure 9, (which is Figure 1 referred to in Example 3) shows a photograph.
 - Figure 10, (which is Figure 2 referred to in Example 3) shows a photograph.
 - Figure 11, (which is Figure 3 referred to in Example 3) shows a photograph.
 - Figure 12, (which is Figure 4 referred to in Example 3) shows a photograph.
- Figure 13, (which is Figure 5 referred to in Example 3) shows a photograph.
 - Figure 14, (which is Figure 6 referred to in Example 3) shows a photograph.
 - Figure 15, (which is Figure 7 referred to in Example 3) shows barcharts.
 - Figure 16, (which is Figure 8 referred to in Example 3) shows a photograph.
 - Figure 17 shows chemical formulae.
- 35 The figures are described more fully in the following example sections.

EXAMPLE 1: STIMULATION OF NEURITE OUTGROWTH

Nerve growth factor acts via retinoic acid synthesis to stimulate neurite outgrowth.

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Nerve growth factor (NGF) stimulates neurite outgrowth from cultured adult dorsal root ganglia (DRG)1. The vitamin A derivative retinoic acid (RA) also induces neurite outgrowth from various embryonic sources, including DRG2,3 Are such similarities in effects of NGF and RA because they are both components of the same genetic cascade leading to neurite outgrowth? RA up-regulates low-and high-affinity NGF receptors3,4 and induces the transcription of NGF itself5, suggesting that RA may be upstream of NGF in the cascade. However, here we show the converse, namely, that NGF is upstream of RA. We show that when adult mouse DRG are cultured in the presence of NGF and a compound that inhibits enzymes involved in RA synthesis, neurite outgrowth does not occur. Conversely, when RA is added along with a blocking antibody to NGF, neurite outgrowth occurs as normal. We further show that NGF induces transcription of both the retinoic acid-synthesizing enzyme RALDH-2 and the retinoic acid receptor- β as well as detectable release of synthesized RA. We propose that RA is required for adult DRG neurite regeneration and that NGF acts upstream of RA to induce its synthesis.

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Cellular effects of RA are mediated by binding to nuclear receptors that are ligand activated transcription factors. There are two classes of receptors, retinoic acid receptors (RARs) and retinoid X receptors (RXRs), with three subtypes of each: α , β and γ 6,7 In addition, there are multiple isoforms of each subtype due to alternative splicing and differential promoter usage. RAR receptors mediate gene expression by forming heterodimers with the RXRs, whereas RXRs can mediate gene expression either as homodimers or by forming heterodimers with orphan receptors such as LXR8 An additional mechanistic association between NGF and RA pathways is suggested by the findings that the nuclear receptor NGFIB heterodimerizes with the RXRs8 and that NGFIB is rapidly induced in PC12 cells by the administration of NGF9.

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Although there is clearly a role for RA in the stimulation of neurite outgrowth from embryonic DRG2,3, it is not yet known if the same occurs in the adult DRG. To test this, we cultured adult mouse DRG in the presence of NGF (100 ng per ml), or RA (100 nM) in delipidated serum for five days. In both cases neurite outgrowth occurred (Fig. 1b). Little or no neurite outgrowth occurred in adult DRG cultured in only delipidated serum (Fig. 1a). Differences in number of neurites were significant (Fig. 2a; 1, 2 and 3). It is important to note that the number of neurites extended from RAor NGF-treated adult DRG, although significantly greater than the number extended from untreated DRG, was smaller than the number obtained using embryonic tissue. When RA was added together with NGF, there was no additive effect of the two treatments (Fig. 1c), and no significant difference was seen between RA, NGF or RA plus NGF groups (Fig. 2a; 2, 3 and 4). Although it may be that both NGF and RA are at individual saturating concentrations, the lack of synergy may also imply that NGF and RA act through the same pathway in order to cause neurite outgrowth. One could imagine either RA inducing the production of NGF5, or NGF inducing the production of RA by stimulating a RA-synthesizing enzyme.

To test which of these hypotheses is most likely, we cultured adult DRG in the presence of NGF and 10 µM disulphiram, a compound which blocks the conversion of retinaldehyde to RA by inhibiting the enzyme aldehyde dehydrogenase10. If RA acts to stimulate NGF production then disulphiram should have no effect on NGF-stimulated neurite outgrowth, whereas if NGF induces RA synthesis then disulphiram should inhibit outgrowth. As shown in Fig. 1d, addition of 10 µM disulphiram along with NGF completely abolished NGF-induced neurite outgrowth (significant difference; Fig. 2a, 2 and 5), whereas addition of DMSO (vehicle for disulphiram) and NGF did not affect neurite outgrowth (Fig. 2a, 2 and 6). To confirm that disulphiram did not affect cell survival within the explants, we performed two types of rescue. In both cases, explants were cultured for eight days in medium supplemented with disulphiram. In the first rescue, 100 nM RA was added to the explants from the beginning of the experiment; in the second, RA was added on day 4. In both cases, significantly greater neurite outgrowth occurred compared to cultures grown in medium supplemented with disulphiram alone (Figs. 1e, f and 2b). These experiments

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also confirm the specificity of disulphiram for the RA synthesis pathway, as RA can rescue the cellular response.

Inhibition of the inductive effect of NGF but not of RA by disulphiram suggests that NGF may precede RA in the cascade leading to neurite outgrowth. To test this, we used a blocking antibody against NGF In the presence of NGF and the blocking antibody, virtually no neurite outgrowth occurred (Fig. 1g; compare to DRG cultured in the presence of NGF alone, Fig. 1b). On the other hand, DRG cultured in the presence of the NGF-blocking antibody and 100 nM RA (Fig. 1h) showed neurite outgrowth equivalent to that obtained with NGF alone (Figs. 1b and 2c).

If NGF is upstream of RA, it should induce synthesis of RA after addition to DRG cultures. To test this prediction, we used an F9 reporter cell line that responds specifically to the presence of RA due to transfection with 1.8 kb of the mouse RAR\$2 gene promoter containing a retinoic acid response element (RARE) linked to the lacZ gene (Sonneveld, E., van den Brink, C. E., van der Leede, B. J., Maden, M. & van der Saag, P. T. (1999) Embryonal carcinoma cell line stably transfected with mRARb2lacZ: sensitive system for measuring levels of active retinoids. Exp.Cell.Res. vol. 250 pp284-297). In the presence of RA, activated cells can be detected after β galactosidase histochemical staining. We first eliminated the possibility that NGF itself activates F9 cells by growing them in the presence of NGF (100 ng per ml), whereupon there was no labeling of the F9 cells above background. We then cultured adult DRG in delipidated serum for five days under three different conditions: in the absence of NGF, in the presence of NGF or in the presence of both NGF and the NGFblocking antibody. Cultured DRG were then sonicated and placed on the F9 reporter cells. NGF-treated DRG homogenates produced a clear RA signal relative to untreated DRG (Fig. 2d). This activation was prevented when the DRG were cultured with blocking antibody in addition to NGF (Fig. 2d).

We next considered which retinoic acid synthesizing enzymes might be induced by NGF. Retinol is converted by a two-step oxidative process to an aldehyde, retinal, which is then oxidized to retinoic acid (for review, see ref. 11). It has been shown that

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retinaldehyde dehydrogenase type 2 (RALDH-2) is expressed in the developing nervous system, including the DRG12. Using RT-PCR, we found strong induction of RALDH-2 by NGF in cultured adult DRG as well (**Fig. 2e**). Lastly, we also found upregulation of the RARβ receptor in NGF-stimulated cultures (**Fig. 2e**), a phenomenon shown to be involved in neurite outgrowth13.

Our results show that RA can stimulate neurite outgrowth from an adult neural tissue, the DRG. NGF similarly stimulates neurite outgrowth from this tissue, and we have demonstrated that it does so by inducing RA synthesis via an enzyme, RALDH-2. In the presence of either a NGF-blocking antibody or an inhibitor of RA synthesis, then NGF fails to act. Thus the most likely sequence of events in the induction of neurite outgrowth by NGF is: NGFRALDH-2RARAR\$\beta\$ neurite outgrowth. We have not yet determined if NGF is directly responsible for inducing RALDH-2, or if some intermediary protein is required for this process. However, as NGFIB is one of the earliest genes induced by NGF9 and its product can heterodimerize with the RXRs8, the NGFIB/RXR heterodimer may be responsible for activating the RALDH-2 gene. Neurotrophins classically have been considered as potential agents for induction of nerve regeneration14 and treatment of neurodegenerative diseases15, but a major problem for their use is lack of effective modes of delivery to the site of the injury. Because RA is required for the regenerative response and it is downstream of NGF, then the problem of delivery to the lesion could be overcome. as RA is a lowmolecular-weight lipophilic compound that can be administered orally. Thus, RA may be of clinical use in neurology.

25 FIGURES FOR EXAMPLE 1

Fig. 1. Neurite outgrowth in adult mouse DRG cultured for five (a-d, g, h) or eight days (e, f) in the presence of delipidated serum plus: (a) no addition; (b) NGF, 100 ng per ml; (c) NGF and 100 nM tRA; (d) NGF and 10 M disulphiram; (e) disulphiram and tRA added on day 0; (f) disulphiram; (g) NGF and blocking antibody (h) NGF-blocking antibody and tRA.

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Fig. 2. (a-c) Neurites produced by adult DRG cultured in cellogen. (a) Effects of NGF, RA and disulphiram at five days (1, no additive: 2, NGF, 100 ng per ml; 3, RA, 100 nM; 4, NGF, 100 ng per ml and RA, 100 nM; 5, 100 ng/ml NGF and 10 M disulphiram: 6, NGF, 100 ng per ml and DMSO). Error bars, s.e.; n = 6, all groups. Differences between NGF-treated (2) and other groups: p < 0.01; p < 0.0001; Student's t-test. (b) RA rescue of DRG treated with 10 M disulphiram (left to right: no RA; 100 nM RA, day 0; 100 nM RA, day 4) Error bars, s.e.; n = 6, all groups. Differences from RA-absent cultures: *p < 0.01, **p < 0.0001; Student's t-test. (c) Effect of NGF-blocking antibody on 5-day DRG cultures. Left, NGF, 100 ng per ml; center, NGF plus blocking antibody; right, blocking antibody plus 100 nM RA; n = 4. Differences from NGF plus blocking antibody: *p < 0.01, **p < 0.0001, Student's ttest. (d) Increase in percentage β-gal-positive F9 cells in response to DRG cultured with or without NGF. Left, no additive; center, NGF, 100 ng per ml; right, NGF with blocking antibody. Differences in percentage β-gal-positive cells from that produced by NGF-treated DRG; *p < 0.025, Student's t-test; for each group, n = 9. (e) RT-PCR analysis of RALDH-2 enzyme and RARB expression in adult DRG cultured with or without NGF (100 ng per ml) for five days. GAPDH was used to indicate presence of cDNA in both samples. Use of F9 reporter cells in studying RA distribution in chick embryo has been described 16.

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EXAMPLE 2: INDUCTION OF NEURITE DEVELOPMENT IN ADULT NEURAL TISSUE.

It is surprisingly shown herein that retinoic acid receptor-β2 induces neurite outgrowth in the adult mouse spinal cord.

Retinoic acid has been shown to be required for neurite outgrowth. We have recently demonstrated that the mechanism of it's action in peripheral nerve regeneration is by activating the retinoic acid receptor $\beta 2$. The adult central nervous system cannot regenerate. Therefore, we have investigated if regenerative failure in the adult spinal cord is related to the expression of retinoic acid receptor $\beta 2$.

Results: We report here that in embryonic mouse spinal cord which can regenerate RARβ2 is up-regulated at concentrations which maximally stimulate neurite outgrowth. In contrast in the adult mouse spinal cord, RARβ2 is not detected nor is it induced by RA and no neurites are extended *in vitro*. When the adult cord is transfected with RARβ2 neurite regeneration can be induced. There is no neurite outgrowth when the cord is transfected with another isoform of RARβ, RARβ4. This shows the importance of receptor specificity in neurite regeneration.

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Conclusion: These data suggest that the loss in regenerative potential of the adult CNS is due in part to the loss of expression of RAR\$\beta\$2 and that it is intrinsic to the neuron itself. We suggest that gene therapy with RAR\$\beta\$2 may result in functional recovery of the injured spinal cord.

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Background The induction of axonal regeneration in the adult central nervous system (CNS) is a major goal in neurobiology. The failure of CNS axons to regenerate under normal circumstances has been attributed to one or a combination of causes: the low abundance of neurotrophic factors; the absence of growth-promoting molecules; the presence of growth-inhibiting molecules. Thus attempts to restimulate axon growth in the CNS have centered on these three possible. When peripheral nerve grafts were used to provide a permissive environment then spinal cord and medulla neurons

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extended axons up to 30mm in the adult rat¹. A similar strategy combined with fibroblast growth factor application resulted in the partial restoration of hind limb function². Neutralisation of neurite growth inhibitors present in myelin with antibodies permitted longer extension of axons than in control young rats³ and led to the recovery of specific reflex and locomotor functions after spinal cord iniurv4. A combination of neurotrophin-3 and these antibodies was successful in inducing long distance regeneration of corticospinal tract (CST) axons⁵. A suspension of olfactory ensheathing cells was also effective in returning locomotor function to the lesioned CST of rats⁶. If neurotrophins act simply to keep axotomised neurons alive⁷ then in these methodologies for inducing regeneration it is the environment surrounding the axons which is the focus of attention rather than the intrinsic capabilities of the neuron itself. However, at least part of the regenerative loss of the CNS is intrinsic to the neuron itself(4 refs). This suggests that the identification of genes that are not expressed in the non regenerating adult CNS but are in the developiong CNS which can regenerate neurites may lead to new stratagies of treatment of spinal cord injuries by gene therapy.

We show here that one such gene is RARβ2 which is activated by retinoic acid (RA) the biologically active metabolite of vitamin A. RA is present in various tissues of the developing embryo and adult animal, especially the nervous system⁸⁻¹³. In its absence, developing neurons of the CNS do not extend neurites into the periphery^{14,15}. Conversely, when applied to cultured neurons, RA induces both a greater number and longer neurites¹⁶ as well as being capable of dictating their direction of growth¹⁷. RA acts at the level of gene transcription because it is a ligand for two classes of nuclear transcription factors, the retinoic acid receptors (RARs) and the retinoid X receptors (RXRs)^{18,19}. There are three members of each class of retinoid receptor a, b and g as well as several isoforms of each member and this diversity may be responsible for the pleitropic effects of RA on cells.

We have been studying the molecular mechanisms of action of RA on neurons and have concluded that one of these retinoic acid receptors, RAR\(\beta\)2 is the crucial transducer of the RA signal in neurons as it is up-regulated in situations where RA stimulates neurite outgrowth²⁰. We hypothesised therefore that the absence or below threshold level of this nuclear receptor in the adult spinal cord may contribute to the failure of this tissue to regenerate axonal projections.

Results and discussion

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Effect of RA on embryonic mouse spinal cord in vitro. 10

We began by confirming that the mouse embryonic spinal cord will respond to RA by extending neurites as do other areas of the embryonic CNS12,17, 21-23 and that this behaviour involves an up-regulation of RAR\$2. E13.5 spinal cord was dissected from mouse embryos placed in a cellogen matrix and cultured in 10 % delipidated serum. All-trans-RA was added at 3 different concentrations (10-8M, 10-7M, 10-6M) and after 5 days the explants were stained with a neurofilament antibody and examined for the presence of neurites. There was an increasing number of neurites emerging from the cultured cord with increasing concentrations of RA with the maximal effect at 10-6M (Fig. 1C, E, G). Even in the absence of RA the embryonic cord extended neurites 20 (Fig. 1A) presumably because of the high endogenous content of RA and its precursor Indeed, when the endogenous synthesis of RA is inhibited with retinol^{9,13}. disulphiram then no neurites are extended²⁴. To demonstrate that the induction of neurites involved the up-regulation of RAR\$2, RT-PCR was performed on cultures after 5 days in the same range of RA treatments. This revealed that RARB2 is 25 normally expressed in embryonic spinal cord at all concentrations of RA used (Fig. 2A, lanes 1-5) and that it is strongly up-regulated after 1 \times 10⁻⁶ M RA treatment (Fig. 2A, lane 5), the same concentration which gives maximal neurite outgrowth.

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Lack of effect of RA on adult mouse spinal cord in vitro.

We next performed an identical series of experiments using 10 month old adult spinal cord rather than the embryonic cord. In contrast to the embryonic cord, RA had no effect on neurite outgrowth at any concentration tested and like the untreated controls, these RA treated adult cords failed to extend any neurites at all (Fig. 1B, D, F, H). Examining the involvement of RAR\$\beta\$2 by RT-PCR revealed that control adult spinal cord had little or no detectable endogenous levels of this receptor (Fig. 2B, lane 1) and that there was no change in its level in response to RA treatment at any concentration (Fig. 2, lanes 2-5), unlike the embryonic cord.

Induction of neurites in adult spinal cord

We therefore hypothesised that it was the lack of RAR\$2 expression which may be responsible for the completely inert behaviour of the adult spinal cord. Our previous observations that adult DRG which do respond to RA by extending neurites also upregulate RAR\$2\$^2\$^4\$ demonstrates that the same behaviour is elicited by embryonic and the appropriate adult neurons and reinforces the differences in regulative behaviour between PNS and CNS neurons. To test our hypothesis we used a defective herpes simplex virus type 1 (HSV-1) vector to transfect pieces of adult (10 months) mouse spinal cord.

Three different transfections were performed, two of which served as controls. Firstly, just the vector containing lacZ (pHSVlacZ); secondly the vector containing RARβ2 (pHSVRARβ2); thirdly the vector containing another isoform of the RARb gene. RARβ4 (pHSVRARβ4). The latter served as a very precise control for transfection since we do not detect the RARβ4 isoform after RA treatment of neurons in our previous experiments²⁰ hence it is not involved in neurite outgrowth. We first ensured that the transfections were successful and that the relevant receptor isoform was expressed in the cultured cord. Pieces of spinal cord were transfected overnight with the appropriate construct and analysed either three or four days later. The

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pHSVlacZ treated cords showed a significant amount of transfection had taken place as judged by b-galactosidase staining of the adult cord (Fig. 3B). RT-PCR demonstrated that transfection with the RAR β 2 vector resulted in the expression of RAR β 2 (Fig. 4, lane 3) but not RAR β 4 (Fig. 4, lane 4) and transfection with the RAR β 4 vector resulted in the expression of RAR β 4 (Fig. 4, lane 8) but not RAR β 2 (Fig. 4, lane 7). In the non transfected cord neither RAR β 2 or RAR β 4 were detected (Fig. 4, lanes 2 and 6).

The effects of these transfections on neurite outgrowth were clear-cut. Transfection with the pHSVlacZ failed to change the behaviour of the cultured adult cord which remained completely un-responsive in terms of neurite outgrowth (Fig. 5A, 12/12 transfections). Similarly, the transfections with pHSVRARβ4 produced no response in the cultured cord which remained inert (Fig. 5C, 12/12 transfections). However, transfections with the pHSVRARβ2 isoform clearly produced a different behaviour and many neurites appeared in the cultures (Fig. 5B, 8/12 transfections). The number of neurites produced in the pHSVRARβ2 cord varivaried between 3 and 23. In the pHSVlacZ transfections there was considerable variability in the number of lacZ-positive cells per explant. This suggests that the variability in neurite number may be due to variability in number of cells transfected.

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These results provide strong support for our hypothesis that the RAR $\beta2$ isoform plays a key role in the induction of neurite outgrowth in response to RA and that this may be a crucial component which fails to be up-regulated in the injured adult CNS. Our hypothesis is based upon several experiments involving either regenerating or non-regenerating neuronal tissues and their response to RA. Thus the embryonic mouse spinal cord, the embryonic mouse DRG and the adult mouse DRG all respond to RA by up-regulating RAR $\beta2$ and extending neurites. In contrast, the adult mouse spinal cord fails to up-regulate RAR $\beta2$ and fails to extend neurites. Furthermore, NGF stimulates neurite outgrowth and acts by up-regulating RAR $\beta2$ and neurite outgrowth from embryonic mouse DRG can be stimulated by a RAR β agonist²⁰.

These results reveal that when the genome of the neuron itself is manipulated then regeneration can be reawakened. This is in contrast to the recent inductions of neurite outgrowth *in vivo* which have concentrated on the inhibitory factors present in the CNS environment 1-5. During development the loss of regenerative capacity of the spinal cord correlates with the appearance of myelin associated neurite growth inhibitory molecules and some of these are thought to be produced by the oligodendrocytes 25. Either the regeneration of neurites we see in our cultures and that seen in cultures where the environment has been manipulated are two different mechanism of neurite regeneration or they are related processes. Support for the latter view is provided by the fact that CNS neurons themselves have been shown to be involved in myelination 26. Therefore it is tempting to speculate that the presence of RARβ2 in neurons during development may regulate genes involved in myelination, and that this process is recapitulated by transfection of the RARβ2 gene in the adult CNS.

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None of the neurites we observed in the RAR β 2 transfected cord elongated over an appreciable distance. This suggests that elongation of the neurite may require the expression of a different set of genes. Evidence that this may be true is shown from regeneration of axons in the adult PNS, where a transition from arborizing to elongating growth depends upon a transcriptional dependent switch²⁷. Alternatively the failure of elongation of neurites in our cultures may be due to the fact that there is likely to be a loss of expression of RAR β 2 over time due to the transient nature of the transfections and that this does not allow enough time for elongation to occur.

Nonetheless we propose that these preliminary data support a role of RARβ2 in the regeneration of neurites in the adult CNS and that gene therapy with this transcript in combination with other treatments may one day lead to functional recovery of the injured spinal cord.

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Methods

Cultures. Spinal cord was dissected from either E13.5 or 10 month old mice and cut into transverse pieces of about 5 mm. These were cultured in cellogen matrix (ICN flow), prepared by mixing 1 volume of 7.5 % sodium bicarbonate, 1 volume of 10x MEM (Gibco) and 8 parts cellogen (ICN flow). The pH was adjusted to 7.5 by dropwise addition of 5M NaOH. Explants were fed every two days. The media consisted of DMEM-F12 with glutamine (Gibco), 6 % glucose, GMS-A (Gibco) 10% delipidated serum and all-*trans*-RA (stock solution, 1 x 10⁻⁵ M, Sigma). On the fifth day they were fixed in 4 % paraformaldyde and stained with the neurofilament antibody, NF200 (Sigma).

RT-PCR analysis. RNA was extracted (trizol, Gibco) and cDNA prepared by the use of a Pharmacia kit as described in the manufactures instructions. The primers used were from GAPDH, RARβ2 and RARβ4, (details upon request). PCR was carried out for 25 cycles for embryonic spinal cord and 40 cycles for adult spinal cord. Amplification was carried out as follows, denaturation for 30 s at 95 °C, annealing for 30 s at 55 °C and extension for 1 min at 72 °C. One fifth of the resultant product was then run on a gel.

Transfections. Virus stocks were prepared and B galactosidase staining carried out as described in ref²⁸. The titres used were: pHSV RAR β 2, 5 x 10⁻⁴ ip/ul, pHSVRAR β 4,, 4 x 10⁻⁴ ip/ul, pHSVlacz 5 x 10⁻⁴ ip/ul.

FIGURES FOR EXAMPLE 2:

Fig.1. Comparison of the effect of retinoic acid on neurite outgrowth on cultured E13.5 (A, C, E, G) and 10 month old adult spinal cord (B, D, F, H). Pieces of spinal cord were cultured in cellogen in the presence of 10 % delipidated serum and RA for a

period of five days. The medium was changed every two days. **A**, **B**, no RA; **C**, **D**, 1 x 10^{-8} M RA; **E**, **F**, 1 x 10^{-7} M RA; **G**, **H**, 1 x 10^{-6} M RA.

Fig.2. Expression of RARβ2 in E13.5 and 10 month old adult spinal cord. Pieces of spinal cord were cultured in the presence of various concentrations of RA for a period of five days after which time RT-PCR analysis of RARβ2 was performed. A. E13.5 (lanes 2-5) and B. 10 month old adult spinal cord (lanes 2-5). Lanes: 1. bluescript/HPA II size markers, 2. no RA, 3. 1 x 10-8 M RA, 4. 1 x 10-7 M RA, 5. 1 x 10-6 M RA. The presence of GAPDH was used to indicate equal amounts of cDNA in the samples.

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<u>Fig.3.</u> Transfection of adult spinal cord with pHSVlacZ. Cultured 10 month old adult spinal cord was transfected with 5×10^{-4} ipu/ul pHSVlacZ overnight and analysed for B galactosidase staining 3 days later. A. non-transfected adult spinal cord. B. adult spinal transfected with pHSVlacZ.

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Fig.4. Transfection of adult spinal cord with either pHSVRARβ2 or pHSVRARβ4. Adult spinal cord was cultured in cellogen and transfected either with 5 x 10⁻⁴ ipu/ul of pHSVRARβ2 or 4 x 10⁻⁴ ipu/ul pHSVRARβ4 overnight. RT-PCR analysis four days after transfection, of RARβ2 (lanes 2-4) and RARβ4 (lanes 6-8) expression in adult spinal cord transfected with Lanes 2, 6 no virus , 3, 7 pHSVRARβ2, 4, 8, pHSVRARβ4. The presence of GAPDH was used to indicate equal amounts of cDNA in the samples. Lanes 1,2 bluescript/HPA II size markers.

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Fig.5. Effect of either pHSVlacZ, pHSVRARβ2 or pHSVRARβ4 transfection in cultured adult spinal cord on neurite outgrowth. Ten month old spinal cord was cultured in cellogen and transfected with either 5 x 10⁻⁴ ipu/ul, pHSVlacZ, 5 x 10⁻⁴ ipu/ul, pHSVRARβ2 or 4 x 10⁻⁴ ipu/ul pHSVRARβ4 overnight, and analysed for neurite staining with NF200 4 days after transfection. Cultured spinal cord transfected with A. pHSVlacZ, B. pHSVRARβ2, C. pHSVRARβ4.

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Fig. 6 is a barchart of the average number of neurites per spinal cord explant.

EXAMPLE 3: NEURITE OUTGROWTH FROM MOUSE GANGLIAL NEURONES

The role of retinoic acid receptors in neurite outgrowth from different populations of embryonic mouse dorsal root ganglia.

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Dorsal root ganglion (DRG) neurons can be categorised into at least three types based upon their neurotrophin requirement for survival. We have analysed the expression of the retinoic acid receptors (RARs) and the retinoid X receptors (RXRs) in NGF, NT-3 and BDNF dependent neurons isolated from embryonic day 13.5 mouse DRG. We show that each population of neurons expressed each of the three RXRs, α , β and γ . However, whilst the NGF and NT-3 dependent neurons expressed each of the RARs α , β and γ the BDNF dependent neurons only expressed RAR α and β . When retinoic acid was added to each of the neuronal classes only the NGF and NT-3 dependent neurons responded by extending neurites, and this response involved the up-regulation of RAR β 2. This specificity was confirmed by the use of receptor selective agonists as only a RAR β 3 selective compound stimulated neurite outgrowth. These results suggest a role for RA acting via RAR β 2 in the outgrowth of neurites.

Introduction

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The neurotrophins are a family of growth factors that are required for the survival of a variety of primary sensory neurons in the developing peripheral nervous system (Snider, 1994). The family includes nerve growth factor (NGF) (Levi-Montalcini, 1987) neurotrophin-3 (NT-3) (Maisonpierre et al., 1990) and brain-derived neurotrophic factor (BDNF) (Barde et al., 1982). They are synthesised in the target fields innervated by peripheral neurons and are thought to be transported by a retrograde mechanism from the target field to support the survival of the developing neurons. The neurotrophins act through receptor tyrosine kinases designated Trk. NGF specifically activates TrkA; BDNF activates TrkB and NT-3 activates TrkC (reviewed in Snider, 1994). Analysis of the phenotypes resulting from loss of function experiments of the neurotrophins and the receptor tyrosine kinases have revealed that

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the dorsal root ganglia (DRG) neurons can be classified into at least three types. Neurons that require NGF for their survival mediate nocioceptive (pain) and thermal receptive functions. In the periphery the axons terminate in the superficial layers of the skin and innervate the superficial laminae of the spinal cord (Crowley et al., 1994; Smeyne et al., 1994). Proprioceptive neurons (sense of position of the limbs in space), which are much larger than NGF type neurons, project into the periphery to the primary endings of muscle spindles and extend a collateral branch to the motor pools in the spinal cord are dependent upon NT-3 for their survival (Ernfors et al., 1994; Farinas et al., 1994; Klein et al., 1994). BDNF neurons are small to medium sized and may include some classes of the mechanoreceptors (Klein et al., 1993: Jones et al., 1994).

In addition to growth factors being involved in the survival of neurons, retinoids can also carry out the same role. Retinoids are a family of molecules derived from vitamin A and include the biologically active metabolite, retinoic acid (RA). The cellular effects of RA are mediated through the action of two classes of receptors, the retinoic acid receptors (RARs) which are activated both by all-trans-RA (tRA) and 9-cis-RA (9-cis-RA), and the retinoid X receptors (RXRs), which are activated only by 9-cis-RA (Kastner et al., 1994; Kleiwer et al., 1994). The receptors are of three major subtypes, α , β and γ , of which there are multiple isoforms due to alternative splicing and differential promoter usage (Leid et al. 1992). The RARs mediate gene expression by forming heterodimers with the RXRs, whilst the RXRs can mediate gene expression as homodimers or by forming heterodimers with a variety of orphan receptors (Mangelsdorf & Evans, 1995). Interestingly, one of the earliest genes induced by NGF in PC12 cells is the orphan receptor NGFI-B (NURR1) (Millbrandt, 1989). This suggests that the growth factor and retinoid mediated pathway in developing neurons can interact. This interaction may be critical for the survival of the neuron because RA has been shown to be involved in the survival and differentiation of neurons (Wuarin and Sidell, 1991; Quinn and De Boni, 1991). Furthermore, many studies on a variety of embryonic neuronal types have shown that RA can stimulate both neurite number and length (reviewed in Maden, 1998) as indeed, can the neurotrophins (Campenot, 1977; Lindsay, 1988; Tuttle and Mathew, 1995). Recently we have shown that RA is

critical for neurite regeneration in adult DRG and that it's synthesis may be regulated by NGF (Corcoran and Maden, 1999).

In the work described here, we use E13.5 mouse DRG to investigate further the nature of the interaction between the retinoid mediated pathway and the growth factor pathway by asking which of the RARs and RXRs are expressed in neurons that are dependent upon different neurotrophins for their survival. We show that a different retinoid receptor profile is indeed expressed in NGF, NT-3 and BDNF neurons and that this profile is altered after an RA treatment which induces neurite outgrowth. Specifically, RAR β 2 is up-regulated in NGF and NT-3 neurons but not in BDNF type neurons. This result was confirmed by the use of receptor selective agonists, as only the RAR β agonist will substitute for RA in inducing neurite outgrowth. These results suggest a role for RA acting via RAR β 2 in the outgrowth of neurites.

15 Materials and Methods

DRG cultures. DRG were obtained from E13.5 mice, freed of non-ganglionic tissue and collected in ice-cold calcium magnesium free phosphate buffered saline. To prepare dissociated cell suspensions the ganglia were treated with 0.05 % trypsin for 15 minutes at 15 °C. The reaction was stopped by the addition of 1% serum and single cells obtained by trituration with a 23 G needle. The cells were then spun at 1000 g for ten minutes and resuspended in media. They were plated out at a density of approximately of 25000 cells/cm² in wells that had been precoated with 100μg/ml poly D lysine for 2 hrs. The cultures were fed every 2 days.

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Culture media consisted of DMEM-F12 with glutamine (Gibco), 6 % glucose, ITS (Gibco). The growth factors used were either 50ng/ml NGF (7s, Promega) 50ng/ml NT3 (Promega) or 50ng/ml BDNF (Promega). Retinoids were used at a concentration of 1 x 10^{-7} M. All-trans-retinoic acid was obtained from Sigma and the receptor agonists were synthesised by CIRD Galderma: CD366 activates RAR α , CD2019 activates RAR β , CD437 activates RAR γ and CD2809 activates all of the RXRs.

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RT-PCR analysis. RNA was extracted (trizol, Gibco) and cDNA prepared by the use of a Pharmacia kit as described in the manufacturer's instructions. The primers used were from mouse RARs. RXRs and GAPDH (details upon request). In order to identify which RAR/RXR receptors were involved neurite outgrowth semi quantitative PCR was used. Amplification was carried out in the linear range for each RAR and RXR and their levels of expression were compared to GAPDH. For the RXRs 28 cycles were performed, while 25 cycles were used for RARα and RARγ and 22 cycles for RARβ and GAPDH. Amplification was carried out as follows, denaturation for 30 s at 95 °C, annealing for 30 s at 55 °C and extension for 1 min at 72 °C. One fifth of the resultant product was then run on a gel and blotted. This was then probed with the appropriate RAR, RXR or GAPDH for normalisation.

In situ Hybridisation: Cells were washed once with PBS and fixed in 4% PFA for 30 mins. They were then washed twice for 5 mins in PBS-0.05% Tween (PBT). Hybridisation was carried out at 55 °C overnight. The buffer consisted of 0.1M Tris-Cl, pH9.5, 0.05M MgCl₂, 0.1M NaCl and 0.1% Tween-20. The cells were then washed sequentially for 15 min. at 65°C in 50 % hybridisation buffer, 50% 2x SSC, 100% 2x SSC, and finally in 0.2% SSC. They were then washed at RT for 5 minutes each in 75% 0.2x SSC, 25% PBT, 50% 0.2x SSC. 50% PBT, 25% 0.2x SSC, 75% PBT, and 100% PBT. The cells were blocked in 2% sheep serum in PBT for 1 hr and incubated with anti DIG antibody overnight at 4°C. The cells were then washed 8 times in PBT for 2 hrs. Colour was developed by using NBT/BCIP according to the manufacturer's instructions (Boehringer Mannheim).

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Immunohistochemistry and measurement of neurite length: Cells were washed once with PBS and fixed in 4% PFA for 30 mins. They were then washed twice for five minutes in PBS-0.05% Tween (PBT). They were then incubated in primary antibody NF200 (sigma) at 4°C overnight and washed 8 times for 2 hrs in PBT. Secondary antibody was then applied for 2 hrs at RT, and the cells again washed 8 times for 2 hrs in PBT. They were then incubated for 5 mins. in PBS containing 0.5

mg/ml DAB and 6% H₂O₂. Neurite length was measured by using NIH image software. The experiments were repeated three times and three random fields were taken for each experiment for analysis. On average there were 40 neurons in each field and the longest neurite branch was measured for a given neuron.

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Results

Expression of receptors by in situ hybridisation.

We first examined the expression of the RARs and the RXRs in primary cultures of E13.5 mouse DRG by in situ hybridisation. Dissociated DRG neurons were cultured in serum free medium either in the presence of NGF, NT-3 or BDNF for a period of five days. In the absence of neurotrophins the cells died. We found that all three types of neurons expressed RXRα (Fig. 1 D, J, P), RXRβ (Fig. 1 E, K, Q) and RXRγ (Fig. 1 F, L, R). In contrast, the RARs showed a differential expression between the three types of neurons. Whilst the NGF and NT-3 dependent neurons expressed RARα (Fig. 1 A, G), RARβ (Fig. 1 B, H) and RARγ (Fig. 1 C, I), the BDNF dependent neurons only expressed RARα (Fig. 1 M) and RARβ (Fig. 1 N). RARγ was not detectable by in situ hybridisation in the BDNF dependent cultures (Fig. 1 O).

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Effect of RA on neurite outgrowth

In order to eliminate any trophic effect of RA on the different populations of neurons we grew the neurons in serum free medium plus the relevant neurotrophin for a period of two days before adding 1 x 10⁻⁷M RA to the cultures for 3 days. Control cultures had no RA added and were maintained in neurotrophin only. There was no significant difference in the numbers of neurons cultured in the presence or absence of RA. This suggests that the effect of RA was on neurite outgrowth and not due to the selective survival of subsets of neurons under the different culture conditions used. In order to analyse neurite outgrowth the cultures were fixed after five days and stained with the monoclonal antibody NF200. Neurite length was measured by NIH image software. The experiment was repeated three times. In total approximately 120 neurons were

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counted in each experiment and the longest neurite was measured from each neuron from which an average neurite length was taken for each treatment. In the absence of RA the BDNF dependent neurons (Fig. 2 E and Fig.7 C column 1) grew neurites whilst the NGF (Fig. 2 A) and NT-3 dependent neurons (Fig. 2 C) showed limited neurite outgrowth (Fig. 7 A and B column 1). In contrast, when RA was added to the medium there was a dramatic increase in the length and number of neurites in the NGF (Fig. 2 B) and NT-3 dependent neurons (Fig. 2 D) and this difference was found to be significant when the length of the neurites were compared (Fig 7 A and B, columns 1 and 2). In contrast RA had no affect on neurite outgrowth of the BDNF dependent neurons (Fig. 2 F and fig. 7 C columns 1 and 2).

Expression of receptors and response to RA by RT-PCR

In order to identify which of the receptors are involved in neurite outgrowth semi-quantitative PCR was carried using primers against the RXRs and the individual RAR isoforms as described in the materials and methods. There was no difference in the expression of the RXRs in each of the three types of neurons cultured with or without RA. In contrast, there were variations in the RAR receptor profiles. Each of the three types of neurons expressed RARα1 (Fig. 3 A, B, C, lane 1) which was strongly upregulated in response to RA in the NGF (Fig. 3 A, lane 8) and NT-3 (Fig. 3 B, lane 8) dependent neurons and only slightly up-regulated in the BDNF dependent neurons (Fig. 3 C, lane 8). It is clear from Fig. 3 that only the RARα1 isoform is readily detectable in these DRG neurons although on over-exposure of the blots the NT-3 dependent neurons expressed the RARα5 and RARα7 isoforms and the BDNF dependent neurons expressed the RARα6 and RARα7 isoforms.

Of the four possible RARβ isoforms only the RARβ2 isoform was detected in all three types of neuron. This isoform was strongly up-regulated by RA in the NGF (Fig. 4 A, lane 6) and NT-3 dependent neurons (Fig. 4 B, lane 6) but not in the BDNF dependent neurons (Fig. 4 C, lane 6) as compared to the non-stimulated cultures (Fig. 4 A, B, C, lane 2)...

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Of the seven RARy isoforms only RARy1 isoform was detected in the neuronal cultures and then only in the NGF (Fig. 5 A, lane 1 and 8) and NT-3 dependent neurons (Fig. 5 B, lane 1 and 8). No RARy1 was detected by RT-PCR in the BDNF dependent neurons.

Receptor selective analogues and neurite outgrowth

The above data suggested that the up-regulation of either RAR α_1 or RAR β_2 may be responsible for the increase of neurite outgrowth observed in the NGF and NT-3 dependent neurons (Fig. 2 B, D). It is more likely to be the RAR β_2 isoform since this receptor is not upregulated in the BDNF dependent neurons and there is no increase in neurite outgrowth when these are stimulated with RA (Fig. 2 F) whereas the RAR α_1 isoform is up-regulated despite a lack of neurite response to RA. In order to distinguish between these two receptors we used receptor selective synthetic retinoids which have been developed specifically to activate individual receptors. CD366 activates RAR α , CD2019 activates RAR β , CD437 activates RAR γ and CD2809 activates all of the RXRs.

In the presence of the RARα agonist there was no significant increase in neurite outgrowth in any neuronal population (Fig. 6 A, E, I and Fig 7 A B and C columns 1 and 3). In contrast, the RARβ agonist significantly increased neurite outgrowth in the NGF and NT-3 dependent neurons compared to non treated neurons (Fig. 6 B, F and Fig 7 A B columns 1 and 4), but did not effect neurite outgrowth in the BDNF dependent neurons (Fig. 6 J and Fig. 7 C columns 1 and 4). When the different neuronal populations were cultured in the presence of the RARγ agonist there was significant decrease in neurite outgrowth in the NGF and NT-3 dependent neurons (Fig. 6 C, G and Fig. 7 A B columns 1 and 5) whereas neurite outgrowth still occurred in the BDNF dependent neurons (Fig. 6 K and Fig. 7 C columns 1 and 5). There was no significant effect on neurite outgrowth in any of the neuronal populations when

they were cultured in the presence of the RXR agonist (Fig. 6 D, H, L and Fig. 7 A B and C columns 1 and 6).

Therefore, in agreement with our RT-PCR data RAR β_2 is required for neurite outgrowth. Furthermore, RAR γ can inhibit neurite outgrowth.

Interrelationships between RARs

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Finally. we attempted to investigate whether there were any regulative interactions between the receptors RAR β 2 and RAR γ 1 since these have opposite affects on neurite outgrowth. In order to examine this we cultured NGF and NT-3 dependent neurons in serum free medium in the presence of either the RAR γ agonist or the RAR β agonist and looked at the levels of receptor expression by semi quantitative RT-PCR 24 hrs. later. The RAR β agonist up-regulated the expression of RAR β 2 in both the NGF and NT-3 dependent neurons (Fig. 8 B, lanes 3 and 6) compared to non-stimulated cultures (Fig.8 lanes 1 and 4) but did not affect the expression of RAR γ 1 (Fig. 7 A, lanes 3 and 6). However, in the presence of the RAR γ agonist, RAR β 2 is reduced in both the NGF and NT-3 dependent neurons (Fig. 8 B, lanes 2 and 5) compared to non-stimulated cultures (Fig. 8 B, lanes 1 and 4). The RAR γ agonist had no effect on the level of RAR γ 1 (Fig. 8 A, lanes 2 and 5). Thus RAR γ 1 can regulate the expression of RAR β 2.

Discussion of Example 3

Our results show that each of the three dorsal root ganglia neuronal populations we have isolated (NGF, NT-3 and BDNF dependent) express both a common set and a unique set of retinoid receptors. With regard to the RXRs they each express RXRα, RXRβ and RXRγ and none of these were found to be directly involved in neurite outgrowth. In contrast, the neurons expressed different RARs depending on the neurotrophin used to select them. The major RAR isoforms that were common to each population were RARα1 and RARβ2. In addition, the NGF and NT-3 populations

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expressed RARy1 which was not expressed in the BDNF population at the time point analysed.

Only the NGF and NT-3 dependent neurons responded to RA by extending neurites whereas the BDNF dependent neurons produced neurites irrespective of the presence or absence of RA. In parallel there was a change in the RAR profile after RA addition. RAR α_1 and RAR β_2 were strongly up-regulated in the NGF and NT-3 dependent neurons whereas in the BDNF dependent neurons only the RAR α_1 was upregulated. This suggested that RAR β_2 was required for the induction of neurite outgrowth and to confirm this observation we used receptor selective agonists.

The development of receptor selective agonists has provided an extremely valuable tool to begin to examine the role of individual receptors in any particular biological process. We showed here that only the RARβ agonist, CD2019, mimiced the effect of RA by inducing neurite outgrowth in NGF and NT-3 neurons thus confirming our RT-PCR results.

We also observed that the RAR γ agonist caused a decrease in neurite outgrowth of the NGF and NT-3 dependent neurons. In an attempt to show whether this was associated with the RAR β 2 expression we examined whether the RAR agonists had any effect on receptor expression. The RAR β agonist upregulated the expression of RAR β 2 but had no effect on the expression of RAR γ 1. In contrast whilst the RAR γ agonist had no effect on the expression of RAR γ 1 it did down-regulate the level of RAR β 2 expression, this phenomenon may also be a prelude to neurite outgrowth. This suggests that the RAR β transcript can be regulated by RAR γ /RXR heterodimers. The lack of increase in neurite outgrowth in response to RA of the BDNF dependent neurons also suggests that in this type of neuron that RAR β may be regulated differently to RAR β in the NGF and the NT-3 dependent neurons at the embryonic stage studied.

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Our results suggest that it is the activation of the RAR pathway that is responsible for neurite outgrowth, since a RXR agonist which activates RXR/orphan receptors had no effect on neurite outgrowth whereas the RAR\$\beta\$ agonist which activates RAR/RXR heterodimers increased the amount of neurite outgrowth. This suggest that if NGF acts via the RXR/orphan receptor pathway by utilising for example NGFI-B then it is not, in these embryonic stages, directly responsible for neurite outgrowth, rather it may be required for neuron survival. Interestingly in the adult the contrast seems to be true. NGF is not required for neuron survival but it is required for neurite outgrowth (Lindsay, 1988). Thus there may be different mechanisms for neurite outgrowth in developing and adult regenerating neurites. However, there is a absolute requirement for RA in neurite outgrowth during development. In the vitamin A deficient quail the neural tube fails to extend neurites into the periphery (Maden et al. 1996; Maden et al., 1998).

The differential response of these neurons to RA and the receptor agonists may have some significance for embryonic and adult tissues which require retinoids for their development and/or survival. In order to activate different RAR/RXR and RXR/orphan receptor combinations there may be different retinoids present in the tissues. Some support for this view is provided by the fact that there are numerous RA generating enzymes which show localised expression during development (McCaffery et al., 1992; Drager & McCaffery, 1995; Godbout et al., 1996: Neiderreither et al., 1997; Ang & Duester, 1997) and each of these enzymes could make different retinoids. Several novel retinoids have so far been discovered, for example 5, 6-epoxyretinoic acid, which is found in the intestine (McCormick et al., 1978), 4-oxoretinol which is the biologically active metabolite that is responsible for the differentiation of murine embryonic F9 cells (Achkar et al., 1996) and 14- hydroxy-4, 14-retroretinol which is found in B lymphocytes (Buck et al., 1991).

Therefore, the embryo may be able to regulate the amount of neurite outgrowth by synthesising different retinoids. By activating RAR β 2/RXR heterodimers neurite outgrowth could occur whereas by activating RAR γ /RXR heterodimers neurite outgrowth could be stopped. In addition the amount of neurite outgrowth could be

regulated by the amount of retinoic acid. For example in the developing mouse spinal cord there are high concentrations of retinoids in the brachial and lumbar enlargements (McCaffery & Drager, 1994). This may be an intrinsic requirement for innervation of the extremities of the limb where the neurites have to travel large distances to reach their final targets, whereas in the thoracic region where the concentration of retinoids are lower such extensive neurite outgrowth would not be required.

Figure Legends for Example 3

Figure 1. Expression of the RARs and RXRs by E13.5 mouse embryo DRG neurons cultured either in the presence of NGF, NT-3 or BDNF. In situ hybridisation of: A-F, NGF neurons; G-L, NT-3 neurons; M-R, BDNF neurons. Expression of: A, G, M, RARα; B, H, N, RARβ; C, I, O, RARγ; D, J, P. RXRα; E, K, , RXRβ; F, L, R, RXRγ.

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- Figure 2. Effect of RA on neurite outgrowth from DRG neurons. DRG neurons were cultured either in the presence of NGF, NT-3 or BDNF for a period of two days at which point 1 x 10⁻⁷ M all-*trans*-RA was added. They were then examined for neurite outgrowth after a total of five days with NF200 antibody. A. NGF; B, NGF + 1 X 10⁻⁷ M RA; C, NT-3; D, NT-3 + 1 X 10⁻⁷; E, BDNF; F, BDNF + 1 X 10⁻⁷ M RA.
- Figure 3. Expression of RARα isoforms in DRG neurons cultured either in the absence or presence of RA. DRG neurons were cultured in the presence of either NGF, NT-3 or BDNF for a period of two days, 1 X 10⁻⁷ M RA was then added and the presence of the RARα isoforms were then assayed by RT-PCR. Controls had no RA added. A, control NGF neurons lanes 1-7; NGF neurons + 1 X 10⁻⁷ M RA lanes 8-14. B, control NT-3 neurons lanes 1-7; NT-3 neurons + 1 X 10⁻⁷ M RA lanes 8-14. C, control BDNF neurons lanes 1-7; BDNF neurons + 1 X 10⁻⁷ M RA lanes 8-14. Lanes: 1 & 8, RARα1; 2 & 9, RARα2; 3 & 10, RARα3; 4 & 11, RARα4; 5 & 12, RARα5; 6

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Figure 4. Expression of RARβ isoforms in DRG neurons cultured either in the absence or presence of RA. DRG neurons were cultured in the presence of either NGF, NT-3 or BDNF for a period of two days, 1 X 10⁻⁷ M RA was then added and the presence of the RARβ isoforms were then assayed by RT-PCR. Controls had no RA added. A. control NGF neurons lanes 1-4; NGF neurons + 1 X 10⁻⁷ M RA lanes 5-8. B, control NT-3 neurons lanes 1-4; NT-3 neurons + 1 X 10⁻⁷ M RA lanes 5-8. C, control BDNF neurons lanes 1-4; BDNF neurons + 1 X 10⁻⁷ M RA lanes 5-8. Lanes: 1 & 5, RARβ1; 2 & 6, RARβ2; 3 & 7, RARβ3; 4 & 8, RARβ4.

Figure 5. Expression of RARy isoforms in DRG neurons cultured either in the absence or presence of RA. DRG neurons were cultured in the presence of either NGF, NT-3 or BDNF for a period of two days, 1 X 10⁻⁷ M RA was then added and the presence of the RARγ isoforms were then assayed by RT-PCR. Controls had no RA added. A, control NGF neurons lanes 1-7; NGF neurons + 1 X 10⁻⁷ M RA lanes 8-14. B, control
NT-3 neurons lanes 1-7; NT-3 neurons + 1 X 10⁻⁷ M RA lanes 8-14. Lanes: 1 & 8, RARγ1; 2 & 9, RARγ2; 3 & 10, RARγ3; 4 & 11. RARγ4; 5 & 12, RARγ5; 6 & 13, RARγ6; 7 & 14, RARγ7.

Figure 6. Effect of RAR and RXR agonists on neurite outgrowth from DRG neurons.

DRG neurons were cultured either in the presence of NGF, NT-3 or BDNF for a period of two days at which point either 1 x 10⁻⁷M of either CD366 (RARα agonist), CD2019 (RARβ agonist), CD437 (RARγ agonist) or CD2809 (pan-RXR agonist) were added to the cultures. Cultures were then stained for neurite outgrowth at five days with the NF200 antibody. A-D, NGF type neurons; E-H, NT-3 type neurons; I-L, BDNF type neurons. Agonists: RARα A, E, I; RARβ B, F, J; RARγ C, G, K; RXR D, H, L.

Figure 7. Effect of retinoid agonists on the length of neurites from A. NGF type neurons; B. NT-3 type neurons, C. BDNF type neurons. Columns 1. no agonist, 2. RA, 3. RAR α , 4. RAR β , 5. RAR γ , 6. RXR. Error bars s.e.m., n = 50. *p < 0.01.

Figure 8. Effect of a RARγ or RARβ agonist on the expression of RARγ1 and RARβ2 expression in DRG neurons cultured in the presence of NGF or NT-3. DRG neurons were cultured in the presence of serum free medium. After two days 1 x 10-7M RARγ or RARβ agonist were then added to the cultures for a period of 24 hrs. RT-PCR analysis of A. RARγ1; B, RARβ2 expression in NGF (lanes, 1-3) and NT-3 (lanes, 4-6) type neurons. Lanes: 1.4, no agonist; 2.5, RARγ agonist; 3, 6, RARβ agonist

SUMMARY

The Examples demonstrate that RAR β 2 and/or an agonist thereof can be used to cause neurite development.

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In particular, we show *inter alia* the use of retinoids to stimulate neurite regeneration in peripheral nerves by activation of RAR β 2.

When peripheral nerves are damaged some regeneration can occur unlike nerves of the central nervous system which show no regeneration. However regeneration of peripheral nerves is limited particularly when there is traumatic nerve injury where there is a loss of nerve tissue such that a gap is created which the regenerating neurite cannot grow across. This delay in nerve regeneration can lead to muscle atrophy and lead to permanent disability.

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In response to peripheral nerve injury neurotrophins are produced. These are a family of growth factors that are required for the survival of a variety of neurons. The family includes nerve growth factor (NGF) neurotrophin-3 (NT-3) and brain-derived neurotrophic factor (BDNF). It was hoped that neurotrophins could be used in the treatment of PNS injuries. However the results have not been encouraging. Two major problems have been encountered, firstly the problem of delivery to the injury, and secondly since different neurons need different neurotrophins a cocktail of them as to be administered in order for all the nerves to regenerate. We have investigated how neurotrophins stimulate neurite regeneration.

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We have found that the vitamin A derivative all-trans-retinoic acid (tRA) like NGF induces neurite outgrowth from various embryonic sources, including PNS. Cellular effects of tRA are mediated by binding to nuclear receptors that are ligand activated transcription factors. There are two classes of receptors, retinoic acid receptors (RARs) and retinoid X receptors (RXRs), with three subtypes of each: α , β and γ . RAR receptors mediate gene expression by forming heterodimers with the RXRs, whereas

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RXRs can mediate gene expression either as homodimers or by forming heterodimers with orphan receptors.

We have found that only RARβ2 is required for neurite outgrowth of all types of neurons we have cultured. Furthermore when adult mouse DRG are cultured in the presence of NGF and an inhibitor of tRA synthesis, neurite outgrowth does not occur. Conversely, when tRA is added along with a blocking antibody to NGF, neurite outgrowth occurs as normal. We have also shown that NGF induces transcription of both the tRA-synthesizing enzyme RALDH-2 and the RARβ2 as well as a detectable release of synthesized tRA.

We propose that the stimulation of RAR $\beta2$ is an intrinsic requirement for the regeneration of neurites in the peripheral nervous system and that crucially this is downstream of the neurotrophins. Therefore in regard to the peripheral nervous system we want to administer retinoids that can activate the RAR $\beta2$ receptor in order for neurite regeneration to occur.

We have extended our observations to the CNS. We have found that the embryonic spinal cord expresses RAR β 2 and that the amount of its expression correlates with the amount of neurite outgrowth. In contrast the adult spinal cord does not express RAR β 2 nor can it regenerate neurites. We have shown that by transfecting RAR β 2 by use of a defective herpes simplex virus type 1 (HSV-1) vector into cultured adult spinal cord we have transformed the normally inert spinal cord into one which can extend neurites. Therefore, we propose that gene therapy of injured spinal cord with RAR β 2 will lead to functional recovery.

The use of retinoid to treat PNS injuries would have at least three major advantages over the use of neurotrophins. Firstly retinoids unlike neurotrophins are small lipophilic molecules which can be easily administered to the site of injury therefore regeneration should occur at a much quicker rate than can be achieved with neurotrophins, this should lead to a reduction in muscle atrophy and consequent

paralysis. Secondly since the stimulation of RARβ2 is crucial to the regeneration of all neurons we have tested only one type of retinoid need be taken circumventing the need to administer a cocktail of neurotrophins. Thirdly retinoids are relatively easy to synthesise unlike neurotrophins.

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Gene therapy with RAR\$2 to treat CNS injuries should lead to functional recovery and therefore the prevention of paralysis.

PNS and CNS injuries occur all over the world unfortunately it is unlikely that the incidence of such injuries will decrease. World wide a 1000 people per million of the population a year suffer spinal cord injury, ten times this number suffer some sort of

PNS injury.

In addition there are three other areas where retinoids would be of use. In leprosy diabetes and AIDS neuropathy occurs (the neurites die) this is equivalent to PNS injury. In both leprosy and diabetes it has been shown that there is a loss of NGF in the skin of both types of patients leading to the loss of pain sensation and inflammation which can lead to ulcer formation. In AIDS patients sensory neuropathy is one of the most common effects of HIV infection, already NGF as been used to treat this condition.

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Hence, we propose that RAR β 2 agonists can be used to treat PNS injuries including neuropathy associated with leprosy, diabetes and AIDS. Gene therapy with RAR β 2 can be used to treat CNS injuries.

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In summation, our results indicate a role for RA acting via RAR β 2 in the outgrowth of neurites from certain classes of neurons.

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The present invention therefore comprises a method of treatment of neurodegenerative disease in which expression of the retinoic acid receptor RAR β 2 is ensured in affected cells or tissues. This may be achieved by treatment with an agonist of the

RAR β 2 receptor or by gene therapy i.e. insertion of the nucleic acid coding for this receptor. The invention may also be seen as the use of these agents in medication for the treatment of peripheral nervous injuries and spinal cord regeneration e.g. in cases of paraplegia.

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All publications mentioned in the specification are herein incorporated by reference. Various modifications and variations of the described methods and system of the present invention will be apparent to those skilled in the art without departing from the scope and spirit of the present invention. Although the present invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in biochemistry, biotechnology, chemistry or related fields are intended to be within the scope of the following claims. For example, it may be possible to substitute some or all of the RARβ2 and/or some or all of the RARβ2 agonist of the present invention with an inhibitor of an antagonist of RARβ2.

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SEQUENCES

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	RARalpha4 forward strand primer 3 ctaagaagacccacacttctg 23
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20	reverse primer 333 gactccacgacatactcagc 314
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RXRbeta

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CLAIMS

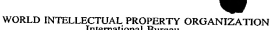
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- 1. Use of RARβ2 and/or an agonist thereof in the preparation of a medicament to cause neurite development.
 - 2. Use of RARβ2 and/or an agonist thereof according to claim 1, wherein said agonist is retinoic acid (RA) and/or CD2019.
- 10 3. Use of RARβ2 and/or an agonist thereof in the preparation of a medicament for the treatment of a neurological disorder.
 - 4. Use of RARβ2 and/or an agonist thereof according to claim 3, wherein said neurological disorder comprises neurological injury.
 - 5. A method of treating a neurological disorder comprising administering a pharmacologically active amount of an RAR β 2 receptor, and/or an agonist thereof.
 - 6. A method according to claim 5, wherein said agonist is RA and/or CD2019.
 - 7. A method according to claim 5 or claim 6, wherein said RARβ2 receptor is administered by an entity comprising a RARβ2 expression system.
- 8. A method of causing neurite development in a subject, said method comprising providing a nucleic acid construct capable of directing the expression of at least part of a RARβ2 receptor, introducing said construct into one or more cells of said subject, and optionally administering a RARβ2 agonist, such as RA and/or CD2019, to said subject.
- 9. A pharmaceutical composition comprising RARβ2 and/or an agonist thereof in admixture with a pharmaceutically acceptable carrier, diluent or excipient; wherein the pharmaceutical composition is for use to cause neurite development.







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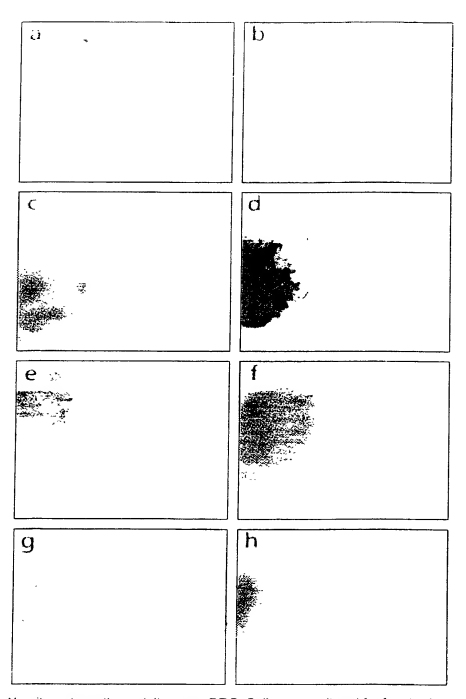
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CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

(54) Title: FACTOR

(57) Abstract

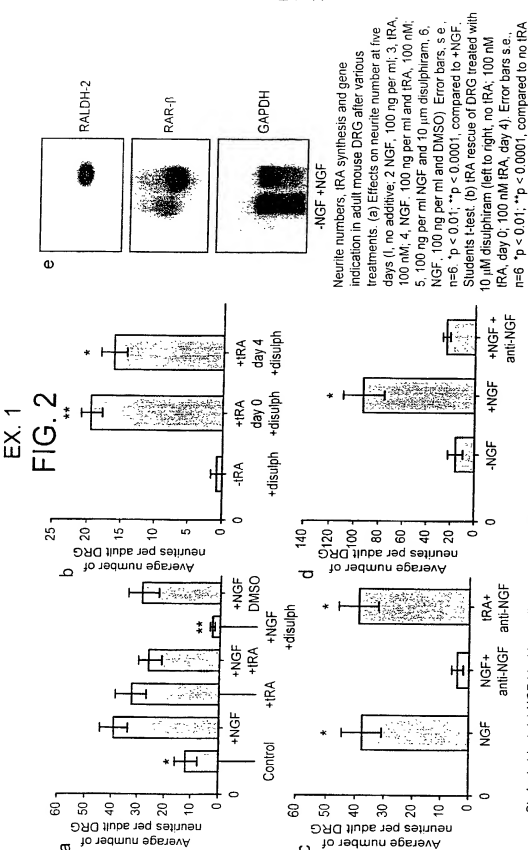
The present invention relates to the use of RAR β 2 and/or an agonist thereof in the preparation of a medicament to cause neurite development.



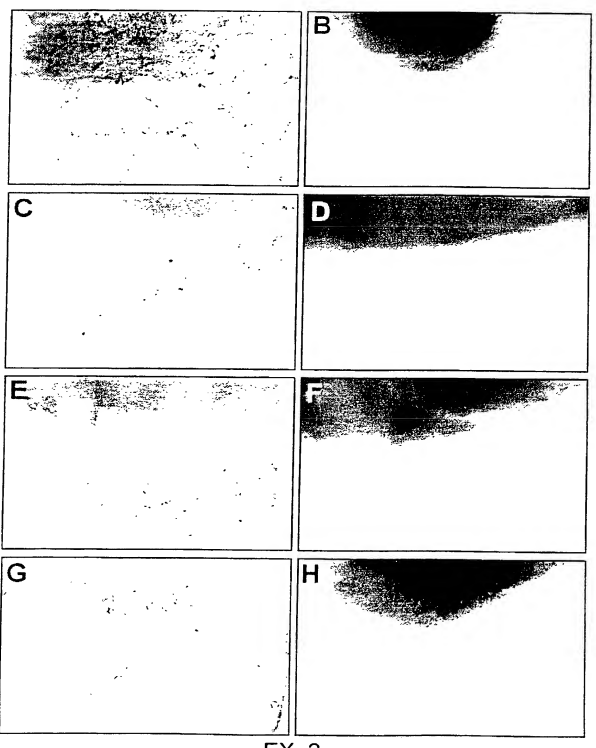
Neurite outgrowth in adult mouse DRG. Cells were cultured for five (a-d, g, h) or eight days (e, f) in the presence of delipidated serum plus (a) no addition. (b) NGF, 100 ng per ml, (c) NGF and 100 nM tRA, (d) NGF and 10 µm disulpiram, (e) disulpiram and tRA added on day 0. (f) disulphiram, (g) NGF and blocking antibody or (h) NGF-blocking antibody and tRA.

EX. 1 FIG. 1

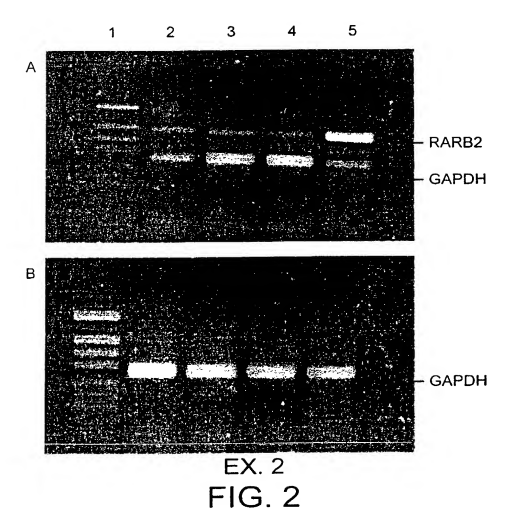




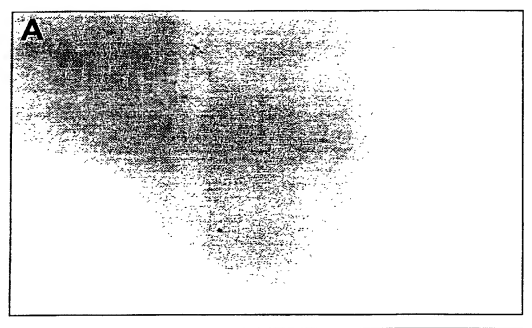
positive F9 cells in response to cultures DRG. Left, no additive; center, NGF 100 ng per ml; right, NGF plus blocking antibody. Cells were counted in three separate fields, and the experiment repeated three times. Error bars s.e., n=9, *p < 0.025 compared to no tNGF, Students t-test. (e) RT-PCR analysis of RALDH-2 enzyme and RARB expression in adult DRG cultured with or without NGF (100 ng per ml) for five days. GAPDH was used to Students t-test. (c) NGF-blocking antibody on 5-day DRG cultures. Left. NGF, 100 ng per ml; center, NGF plus blocking antibody; right, blocking antibody plus 100nM tRA. Error bars s.e., n=6. *p < 0.01, comared to NGF+ α-NGF, Students t-test. (d) Increase in percentage β-galactosidase ndicate presence of cDNA in both samples

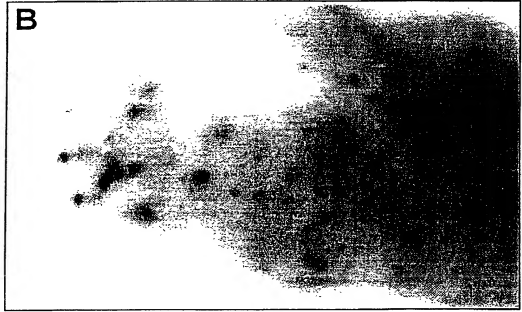


EX. 2
FIG. 1
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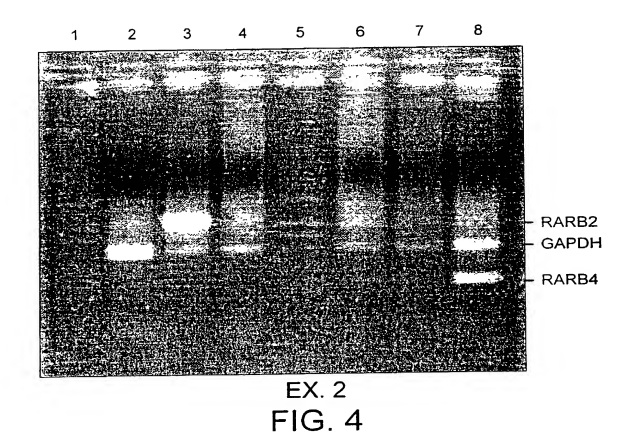






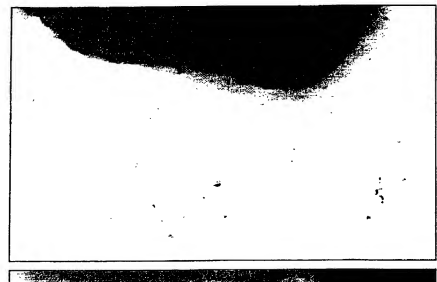


EX. 2 FIG. 3

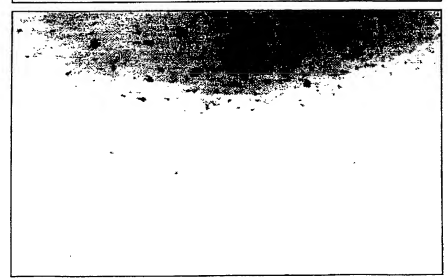


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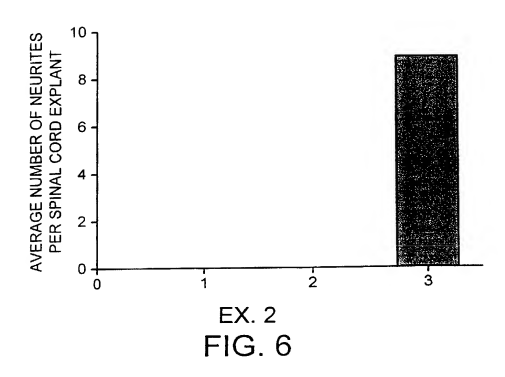
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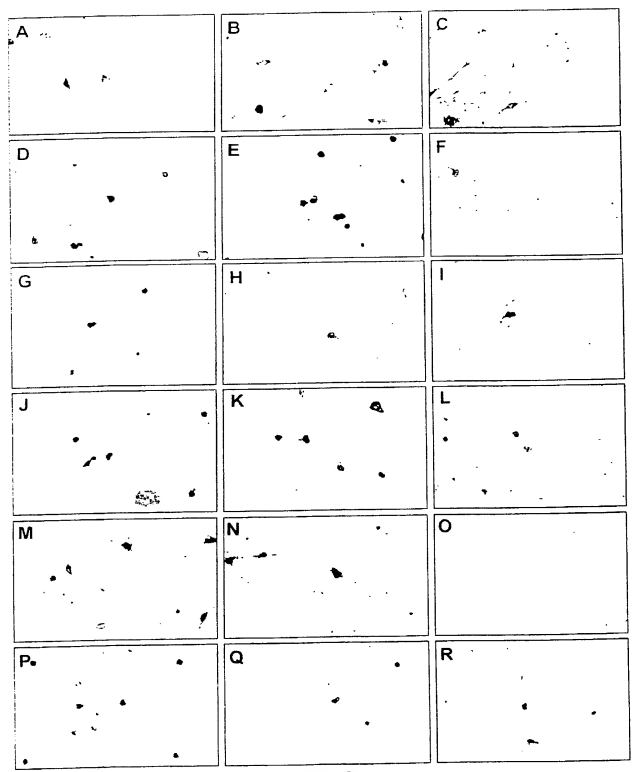




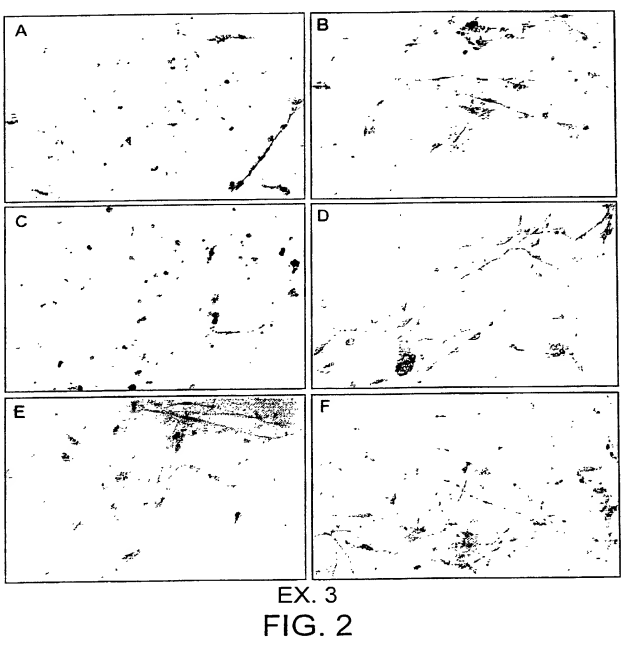


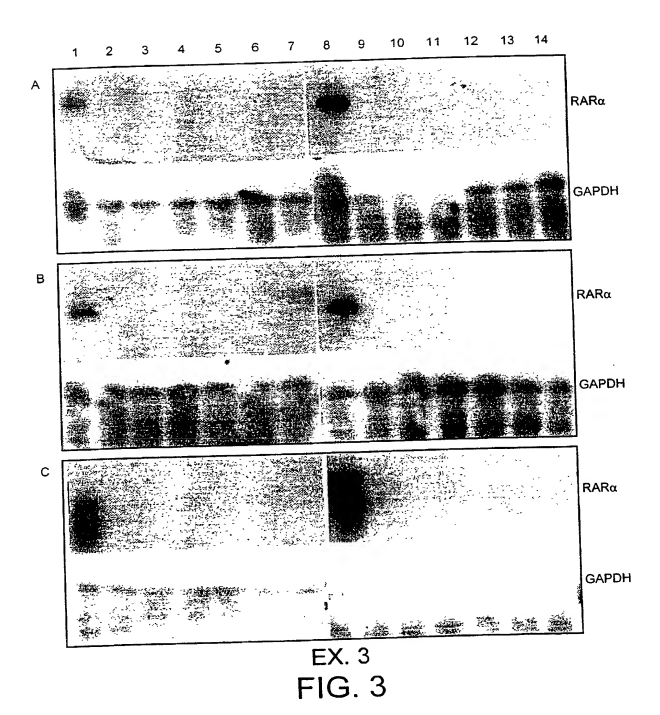
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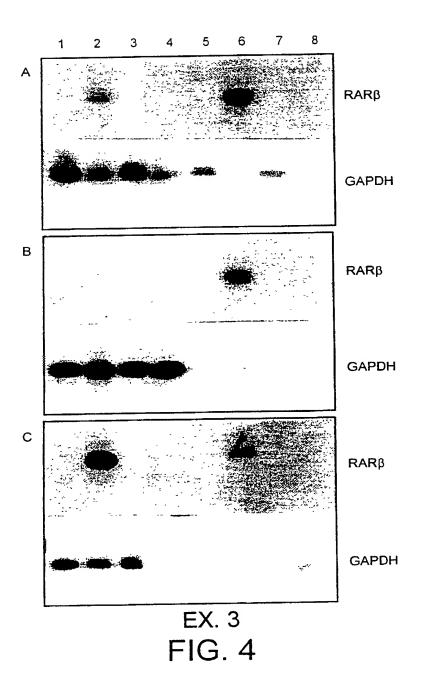


EX. 3 FIG. 1



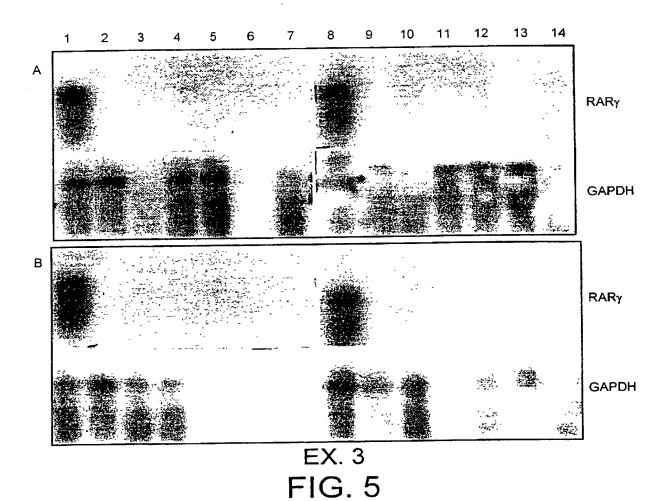


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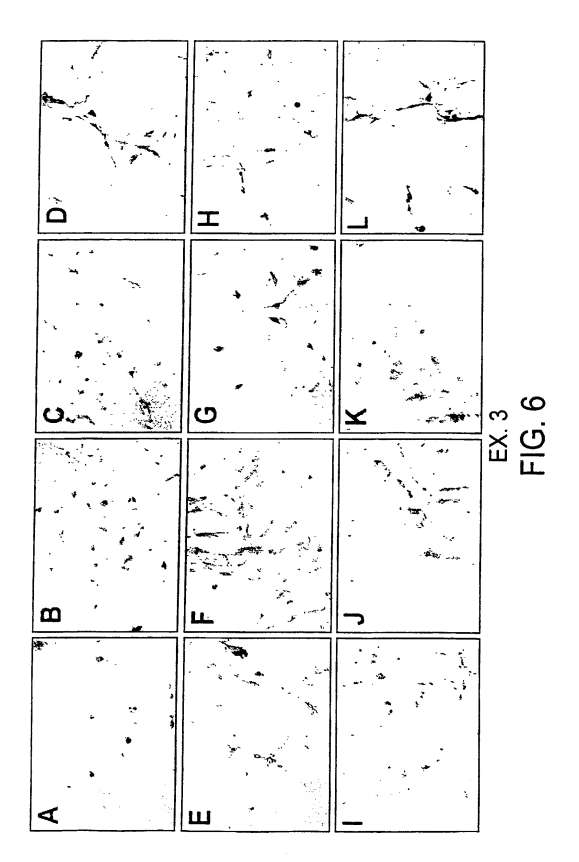


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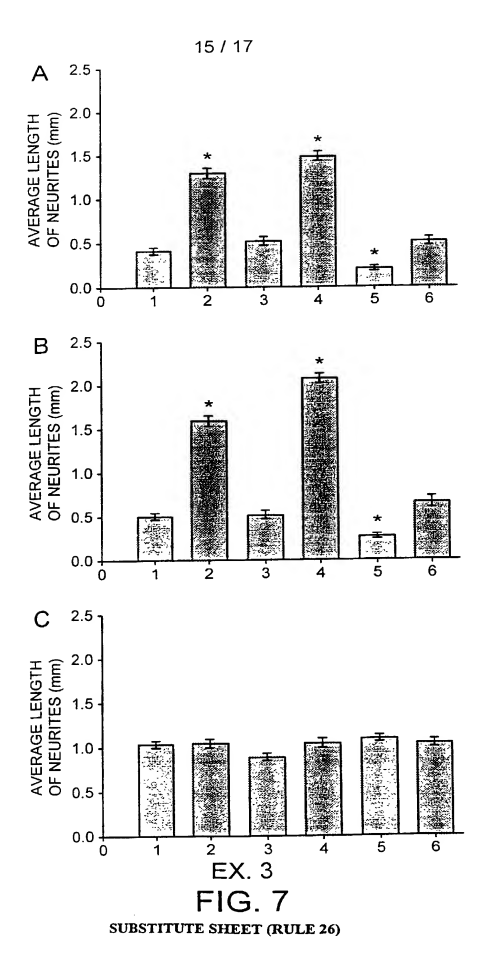
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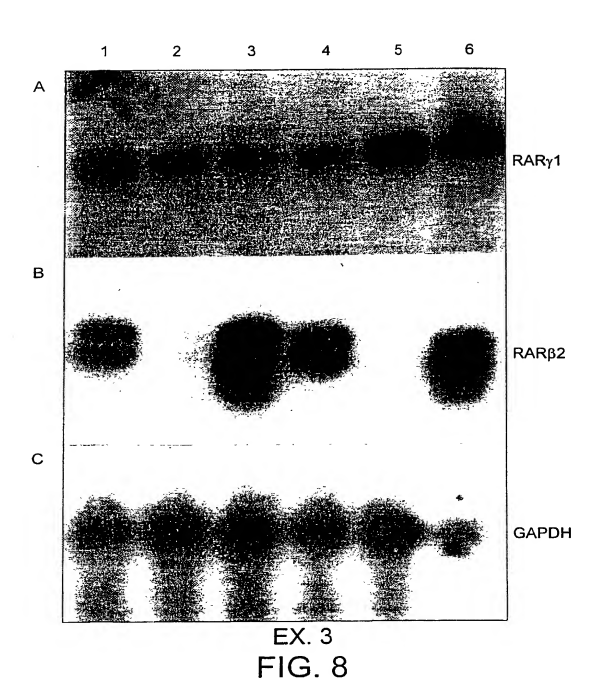


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550-266 P008690US CTH WJR LSB Nixon & Vanderhye P.C. (10/99) (Domestic Non-Assigned/Foreign) Page 1

RULE 63 (37 C.F.R. 1.63) INVENTORS DECLARATION FOR PATENT APPLICATION IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

As a below named inventor, I hereby declare that my residence, mailing address and citizenship are as stated below next to my name, and I believe I am the original, first and sole inventor (if only one name is listed below) or an original, first and joint inventor (if plural names are listed below) of the subject matter which is claimed and for which a patent is sought on the invention entitled: FACTOR the specification of which (check applicable box(s)): is attached hereto (Atty Dkt. No. 550-266) as U.S. Application Serial No. was filed on was filed as PCT International application No. PCT/GB00/01211 30 March 2000 \boxtimes and (if applicable to U.S. or PCT application) was amended on I hereby state that I have reviewed and understand the contents of the above identified specification, including the claims, as amended by any amendment referred to above. I acknowledge the duty to disclose to the Patent Office all information known to me to be material to patentability as defined in 37 C.F.R. 1.56. I hereby claim foreign priority benefits under 35 U.S.C. 119/365 of any foreign application(s) for patent or inventor's certificate listed below and have also identified below any foreign application for patent or inventor's certificate having a filing date before that of the application on ch priority is claimed or, if no priority is claimed, before the filing date of this application: rity Foreign Application(s): Day/Month/Year Filed **Application Number** Country 31 03 1999 9907461.9 Great Britain I hereby claim the benefit under 35 U.S.C. §119(e) of any United States provisional application(s) listed below. Date/Month/Year Filed **Application Number** I hereby claim the benefit under 35 U.S.C. 120/365 of all prior United States and PCT international applications listed above or below: Status: patented Prior U.S./PCT Application(s): pending, abandoned Application Serial No. Day/Month/Year Filed PCT/GB00/01211 30 March 2000 Pending I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patent issued thereon. And on behalf of the owner(s) hereof, I hereby appoint NIXON & VANDERHYE P.C., 1100 North Glebe Rd., 8th Floor, Arlington, VA 22201-4714, telephone number (703) 815-4000 (to whom all applications). Floor, Arlington, VA 22201-4714, telephone number (703) 816-4000 (to whom all communications are to be directed), and the following attorneys thereof (of the same address) individually and collectively owner's/owners' attorneys to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith and with the resulting patent: Larry S. Nixon, <u>25640</u>; Arthur R. Crawford, <u>25327</u>; James T. Hosmer, <u>30184</u>; Robert W. Faris, <u>31352</u>; Richard G. Besha, <u>22770</u>; Mark E. Nusbaum, <u>32348</u>; Michael J. Keenan, <u>32106</u>; Bryan H. Davidson, <u>30251</u>; Stanley C. Spooner, 27393; Leonard C. Mitchard, 29009; Duane M. Byers, 33363; Jeffry H. Nelson, 30481; John R. Lastova, 33149; H. Warren Burnam, 29366; Mary J. Wilson, 32955; J. Scott Davidson, 33489; Alan M. Kagen, 36178; Robert A. Molan, 29834; B. J. Sadoff, 3663; James D. Berquist, 34776; Updeep S. Gill, 37334; Michael J. Shea, 34725; Donald L. Jackson, 41090; Michelle N. Lester, 32331; Frank P. Presta, 19828; Joseph S. Presta, 35329; Joseph A. Rhoa, 37515; Raymond Y. Mah, 41426; Chris Comuntzis, 31097; Gary T. Tanigawa, 43180. I also authorize Nixon & Vanderhye to delete any attorney names/numbers no longer with the firm and/o act and rely solely on instructions directly communicated from the person, assignee, attorney, firm, or other organization sending instructions to flux and entry on behalf of the owner(s). Inventor's Signature: British Malcolm Maden Inventor: (citizenship) (last) (state/country) England Residence: (city) Biomedical Sciences Division/The Randall Institute, Developmental Biology Reearch Centre, King's College Mailing Address: London, 26/29 Drury Lane, London, England (Zip Code) WC2B 5RL Inventor's Signature: British inventor: Jonathan P.T. Corcoran (citizenship) (last) (first) (state/country) England Residence: (city) London Biomedical Sciences Division/The Randall Institute, Developmental Biology Research Centre, King's College Mailing Address: London, 26/29 Drury Lane, London, England WC2B 5RL (Zip Code)

See attached sheet(s) for additional inventor(s) information!!